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201-16472B

IUCLID

Data Set

Existing Chemical : 1D: 6863-58-7 **CAS No.** : 6863-58-7

EINECS Name : 2,2'-oxybisbutane

EC No. : 229-961-6 Common name : sec-Butyl Ether

Molecular Formula : C8H18O

Producer related part

Company : ExxonMobil Biomedical Sciences Inc.

Creation date : 16.10.2006

Substance related part

Company : ExxonMobil Biomedical Sciences Inc.

Creation date : 16.10.2006

Status :

Memo : ExxonMobil Chemical Company - HPV

Printing date : 14.11.2006
Revision date :

Date of last update : 14.11.2006

Number of pages : 31

Chapter (profile) : Chapter: 1, 2, 3, 4, 5, 6, 7, 8, 10
Reliability (profile) : Reliability: without reliability, 1, 2, 3, 4

Flags (profile) : Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE),

Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. General Information

ld 6863-58-7 **Date** 14.11.2006

Molecular formula : C8H18O1 Molecular weight : 130.23

Petrol class

07.11.2006

Purity type : typical for marketed substance

Substance type : organic

Physical status
Purity
Colour
Odour

08.11.2006

bis (2-butyl) ether

07.11.2006

di-sec-Butyl Ether

07.11.2006

sec-Butyl Ether

07.11.2006

1. Ge	eneral Information		6863-58-7 14.11.2006
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1. General Information

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1.10 SOURCE OF EXPOSURE

Type of search

: Internal and External

Chapters covered

:

Date of search

: 16.10.2006

Remark : Search covered all P

Search covered all Physical Chemical Properties, Environmental Fate, Aquatic and Mammalian Toxicity endpoints related to the CAS number.

08.11.2006

Type of search

Internal and External

Chapters covered

:

Date of search

: 25.05.2005

Remark: Search covered all Physical Chemical Properties, Environmental Fate,

Aquatic and Mammalian Toxicity endpoints related to the CAS number.

08.11.2006

Type of search

Internal and External

Chapters covered Date of search

: 20.05.2005

Remark: Search covered all Physical Chemical Properties, Environmental Fate,

Aquatic and Mammalian Toxicity endpoints related to the CAS number.

08.11.2006

ld 6863-58-7 **Date** 14.11.2006

Value : =-100 °C

Sublimation

Method : other: measured

Year

GLP : no data

Test substance : other TS: CAS No. 6863-58-7; sec-butyl ether

Test substance : CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

Although the original data were not retrieved and reviewed for quality, they

were developed following acceptable test methods and therefore

considered reliable. Value was provided by the experimental database of

the EPIWIN program.

Flag : Critical study for SIDS endpoint

14.11.2006 (6)

Value : = -73 °C

Sublimation

Method : other: calculated

Year : 2003 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Method : Calculated values using MPBPWIN version 1.41, a subroutine of the

computer program EPIWIN version 3.12

Test condition: Melting Point estimations performed by MPBPWIN are based on the

average result of the calculation methods of K. Joback and Gold and Ogle.

Joback's Method is described in Joback, K.G. 1982. A Unified Approach to Physical Property Estimation Using Multivariate Statistical Techniques. In The Properties of Gases and Liquids. Fourth Edition. 1987. R.C. Reid, J.M.

Prausnitz and B.E. Poling, Eds.

The Gold and Ogle Method simply uses the formula

Tm = 0.5839Tb, where Tm is the melting point in Kelvin and Tb is the

boiling point in Kelvin.

Test substance: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

The result is a calculated value based on the chemical structure and

represents a potential melting point for the substance with the CAS number

listed under test substance.

14.11.2006 (6)

Value : = 116 °C at 1013 hPa

Decomposition

Method : other: calculated

Year : 2003 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition : Boiling point calculated by MPBPWIN subroutine, which is based on the

method of S. Stein and R. Brown in "Estimation of Normal Boiling Points from Group Contributions". 1994. J. Chem. Inf. Comput. Sci. 34: 581-587.

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Test substance: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

The result is a calculated value based on the chemical structure and represents a potential boiling point for the substance with the CAS number

listed under test substance.

Flag : Critical study for SIDS endpoint

14.11.2006 (6)

Type :

Value : = .759 g/cm³ at 25 °C Method : other: measured

Year : 1935 GLP : no data

Test substance : other TS: CAS No. 6863-58-7; sec-butyl ether

Test substance : CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

Although the original data were not retrieved and reviewed for quality, they were reported in the peer-reviewed literature and therefore considered

reliable.

Flag : Critical study for SIDS endpoint

14.11.2006 (3)

Value : = 21.7 hPa at 25 °C

Decomposition :

Method : other (measured)

Year : 1994 GLP : no data

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test substance: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

Although the original data were not retrieved and reviewed for quality, they

were developed following acceptable test methods and therefore

considered reliable. Value was provided by the experimental database of

the EPIWIN program.

Reference given in EPI Suite: Yaws, CL (1994B).

Flag : Critical study for SIDS endpoint

14.11.2006 (6)

Value : = 29.7 hPa at 25 °C

Decomposition

Method : other (calculated)

Year : 2003 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Method : Vapor pressure calculation by MPBPWIN ver. 1.40 using calculation

method of Grain.

Remark : EPIWIN is used and advocated by the US EPA for chemical property

ld 6863-58-7 **Date** 14.11.2006

estimation.

Test substance : CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

The result is a calculated value based on the chemical structure and represents a potential vapor pressure for the substance with the CAS

number listed under test substance.

14.11.2006 (6)

Partition coefficient : octanol-water Log pow : = 3.35 at 25 °C

pH value

Method : other (measured)

Year : 1992 GLP : no data

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition: Specific methods were not given in the document reporting the measured

value. Measurements were performed at the Chemicals Inspection and

Testing Institute of Japan.

Test substance : CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

Although the original data were not retrieved and reviewed for quality, they were developed at a well respected testing facility and therefore considered

reliable.

Flag : Critical study for SIDS endpoint

14.11.2006 (2)

Partition coefficient : octanol-water Log pow : = 2.87 at 25 °C

pH value

Method : other (calculated)

Year : 2003 GLP : no

Test substance : other TS: CAS No. 6863-58-7; sec-butyl ether

Method : Calculated values using KOWWIN version 1.67, a subroutine of the

computer program EPIWIN version 3.12

Test condition : Octanol / Water Partition Coefficient estimations performed by KOWWIN

are based on an atom/fragment contribution method of W. Meylan and P. Howard in "Atom/fragment contribution method for estimating octanol-water

partition coefficients". 1995. J. Pharm. Sci. 84:83-92.

Test substance: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

The result is a calculated value based on the chemical structure and represents a potential partition coefficient for the substance with the CAS

number listed under test substance.

14.11.2006 (6)

Solubility in : Water

Value : = 330 mg/l at 25 °C

pH value

concentration : at °C

Temperature effects : Examine different pol. :

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pKa : at 25 °C

Description : Stable : Deg. product :

Method : other: measured

Year : 1992 GLP : no data

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition: Specific methods were not given in the document reporting the measured

value. Measurements were performed at the Chemicals Inspection and

Testing Institute of Japan.

Test substance: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

Although the original data were not retrieved and reviewed for quality, they were developed at a well respected testing facility and therefore considered

reliable.

Flag : Critical study for SIDS endpoint

14.11.2006 (2)

Solubility in : Water

Value : = 327 mg/l at 25 °C

pH value

concentration : at °C

Temperature effects

Examine different pol.

pKa : at 25 °C

Description :

Stable : Deg. product :

Method : other: calculated

Year : 2003 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Method: Calculated values using WSKOWWIN version 1.41, a subroutine of the

computer program EPIWIN version 3.12

Test condition: Water Solubility estimations performed by WSKOWWIN are based on a

Kow correlation method described by W. Meylan, P. Howard and R. Boethling in "Improved method for estimating water solubility from octanol/water partition coefficient". Environ. Toxicol. Chem. 15:100-106.

1995.

Test substance : CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

The result is a calculated value based on the chemical structure and represents a potential water solubility for the substance with the CAS

number listed under test substance.

14.11.2006 (6)

2. PI	hysico-Chemical Data		6863-58-7 14.11.2006
2.9	FLAMMABILITY		
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Type : water

Light source :

Light spectrum: nm

Relative intensity: based on intensity of sunlight

Deg. product

Method : other (calculated): Technical discussion

Year : 2006

GLP

Test substance : other TS: CAS No. 6863-58-7; sec-butyl ether

Result: Photolysis as a Function of Molecular Structure

The direct photolysis of an organic molecule occurs when it absorbs sufficient light energy to result in a structural transformation (Harris, 1982). The reaction process is initiated when light energy in a specific wavelength range elevates a molecule to an electronically excited state. However, the excited state is competitive with various deactivation processes that can result in the return of the molecule to a non excited state.

The absorption of light in the ultra violet (UV)-visible range, 110-750 nm, can result in the electronic excitation of an organic molecule. Light in this range contains energy of the same order of magnitude as covalent bond dissociation energies (Harris, 1982). Higher wavelengths (e.g. infrared) result only in vibrational and rotational transitions, which do not tend to produce structural changes to a molecule.

The stratospheric ozone layer prevents UV light of less than 290 nm from reaching the earth's surface. Therefore, only light at wavelengths between 290 and 750 nm can result in photochemical transformations in the environment (Harris, 1982). Although the absorption of UV light in the 290-750 nm range is necessary, it is not always sufficient for a chemical to undergo photochemical degradation. Energy may be re-emitted from an excited molecule by mechanisms other than chemical transformation, resulting in no change to the parent molecule.

A conservative approach to estimating a photochemical degradation rate is to assume that degradation will occur in proportion to the amount of light wavelengths >290 nm absorbed by the molecule (Zepp and Cline, 1977).

Saturated hydrocarbons and R-OH groups do not absorb light above 290 nm (Harris J, 1982a). Therefore, these moieties are stable in regard to direct photolytic processes. Ethers are also stable as this group absorbs UV light in the far UV region, below 220 nm (Mill T, 2000), therefore, direct photolysis will not be an important transformation process for the degradation of sec-butyl ether in the environment.

References:

Harris, J. C. 1982. "Rate of Aqueous Photolysis," Chapter 8 in: W. J. Lyman, W. F. Reehl, and D. H. Rosenblatt, eds., Handbook of Chemical Property Estimation Methods, McGraw-Hill Book Company, New York, USA.

Zepp, R. G. and D. M. Cline. 1977. Rates of Direct Photolysis in the Aqueous Environment, Environ. Sci. Technol., 11:359-366.

Boethling, R.S., Mackay, D. 2000. Handbook of Property Estimation

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Methods for Chemicals, CRC Press, Boca Raton, FL, USA.

Test substance : CAS No. 6863-58-7; sec-butyl ether **Flag** : Critical study for SIDS endpoint

14.11.2006 (5)

Type : air
Light source : Sun light
Light spectrum : nm

Relative intensity : based on intensity of sunlight

INDIRECT PHOTOLYSIS

Sensitizer : OH

Conc. of sensitizer : 1500000 molecule/cm³

Rate constant : = $.00000000000320784 \text{ cm}^3/(\text{molecule*sec})$

Degradation: % after

Deg. product

Method : other (calculated): Calculated values using AOPWIN version 1.91, a

subroutine of the computer program EPIWIN version 3.12

Year : 2003 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Result: Atmospheric Oxidation Potential

In the environment, organic chemicals emitted into the troposphere are degraded by several important transformation processes. The dominant transformation process for most compounds is the daylight reaction with hydroxyl (OH-) radicals (Atkinson, 1988, 1989). The rate at which an organic compound reacts with OH- radicals is a direct measure of its atmospheric persistence (Meylan and Howard, 1993).

AOPWIN estimates the rate constant for the atmospheric, gas-phase reaction between photochemically produced hydroxyl radicals and organic chemicals. The rate constants estimated by the program are then used to calculate atmospheric half-lives for organic compounds based upon average atmospheric concentrations of hydroxyl radicals.

Since the reactions only take place in the presence of sunlight, the atmospheric half-lives are normalized for a 12-hour day.

Calculated* OH- Rate Constant half-life (days) (cm3/molecule-sec)

0.333 32.0784 E-12

References:

Atkinson, R. 1988. Estimation of gas-phase hydroxyl radical rate constants for organic chemicals. Environ. Toxicol. Chem. 7:435-442.

Atkinson, R. 1989. Kinetics and mechanisms of the gas-phase reactions of the hydroxyl radical with organic compounds. J. Phys. Chem. Ref. Data Monograph No. 1, Amer. Inst. Physics & Amer. Chem. Soc., NY.

Meylan, W.M. and P.H. Howard. 1993. Computer estimation of the atmospheric gas-phase reaction rate of organic compounds with hydroxyl radicals and ozone. Chemosphere 12:2293-2299.

Test condition

Indirect photodegradation, or atmospheric oxidation potential, is based on the structure-activity relationship methods developed by R. Atkinson.

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(6)

Temperature: 25°C Sensitizer: OH radical

Concentration of Sensitizer: 1.5 E6 OH radicals/cm3

Test substance: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

The results include calculated data based on chemical structure as

modeled by AOPWIN. The data represent a potential atmospheric half-life

range for the test substance.
Critical study for SIDS endpoint

Flag : Critical study for SIDS endpoint 14.11.2006

 Type
 : abiotic

 t1/2 pH4
 : at °C

 t1/2 pH7
 : at °C

 t1/2 pH9
 : at °C

Deg. product

Method : other: technical discussion

Year : 2006

GLP

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Result : Hydrolysis as a Function of Molecular Structure

Hydrolysis of an organic molecule occurs when a molecule (R-X) reacts with water (H2O) to form a new carbon-oxygen bond after the carbon-X bond is cleaved (Gould, 1959; Harris, 1982). Mechanistically, this reaction is referred to as a nucleophilic substitution reaction, where X is the leaving group being replaced by the incoming nucleophilic oxygen from the water molecule.

Chemicals that are susecptible to hydrolysis contain functional groups that can be displaced by a nucleophilic substitution reaction. Substances that have a potential to hydrolyze include alkyl halides, amides, carbamates, carboxylic acid esters and lactones, epoxides, phosphate esters, and sulfonic acid esters (Neely, 1985). The lack of a suitable leaving group renders a compound resistant to hydrolysis.

Sec-butyl ether is expected to be resistant to hydrolysis because it lacks a functional group that is hydrolytically reactive (Harris, 1982b). Therefore, hydrolysis will not contribute to its removal from the environment.

References:

Gould, E.S. (1959), Mechanism and Structure in Organic Chemistry, Holt, Reinhart and Winston, New York, NY, USA.

Harris, J.C. (1982), "Rate of Hydrolysis," Chapter 7 in: W.J. Lyman, W.F. Reehl, and D.H. Rosenblatt, eds., Handbook of Chemical Property Estimation Methods, McGraw-Hill Book Company, New York, NY, USA.

Neely, W. B. 1985. Hydrolysis. In: W. B. Neely and G. E. Blau, eds. Environmental Exposure from Chemicals. Vol I., pp. 157-173. CRC Press, Boca Raton. FL. USA.

Test substance : CAS No. 6863-58-7; sec-butyl ether **Flag** : Critical study for SIDS endpoint

14.11.2006 (4)

ld 6863-58-7 **Date** 14.11.2006

3.1.3 STABILITY IN SOIL

Type : fugacity model level III

Media : other: air - sediment(s) - soil - water

Air : % (Fugacity Model Level I)

Water : % (Fugacity Model Level I)

Soil : % (Fugacity Model Level I)

Biota : % (Fugacity Model Level II/III)

Soil : % (Fugacity Model Level II/III)

Method : other: Calculation according Mackay, Level III

Year : 2006

Method : The EQC Level III model is a steady state model that is useful for

determining how the medium of release affects environmental fate. Level III fugacity allows non-equilibrium conditions to exist between connected media as steady state, and illustrate important transport and transformation

processes.

Physicochemical input values for the model were calculated using the EPIWIN Estimation v 3.12 program. Measured input values were also used where available and obtained from the EPIWIN database or other reliable sources. Distribution data from the equilibrium model provide basic information on the potential partitioning behavior of chemicals between selected environmental compartments (i.e., air, water, soil, and sediment).

Input values used:

Molecular mass = 130.23 g/mol Water solubility = 327 mg/L Vapour pressure = 2170 Pa

log Kow = 2.87

Melting point = -100 deg C

Degradation half-lives:

Air - 4.0 hrs Water - 360 hrs Soil - 7200 hrs Sediment - 72000 hrs

This model was run assuming the default emissions.

Result : Air - 3.2%

Water - 44.8% Soil - 51.1% Sediment - 0.9%

Test substance: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated and not measured.

Flag : Critical study for SIDS endpoint

14.11.2006 (8)

ld 6863-58-7 **Date** 14.11.2006

Type : fugacity model level I

Media : other: air - biota - sediment(s) - soil - water

Air : % (Fugacity Model Level I)
Water : % (Fugacity Model Level I)
Soil : % (Fugacity Model Level I)
Biota : % (Fugacity Model Level II/III)
Soil : % (Fugacity Model Level II/III)

Method : other: Calculation according Mackay, Level I

Year : 2006

Method : The EQC Level I is a steady state, equilibrium model that utilizes the input

of basic chemical properties including molecular weight, vapor pressure, and water solubility to calculate distribution within a standardized regional

environment.

Physicochemical input values for the model were calculated using the EPIWIN Estimation v 3.12 program. Measured input values were also used where available and obtained from the EPIWIN database or other reliable sources. Distribution data from the equilibrium model provide basic information on the potential partitioning behavior of chemicals between selected environmental compartments (i.e., air, water, soil, and sediment).

Input values used:

Molecular mass = 130.23 g/mol Water solubility = 327 mg/L Vapour pressure = 2170 Pa

log Kow = 2.87

Melting point = -100 deg C

Result : Air - 99.1%

Water - 0.5% Soil - 0.4%

Sediment - <0.01% Suspended Sed - <0.01%

Biota - < 0.01%

Test substance : CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated and not measured.

Flag : Critical study for SIDS endpoint

14.11.2006 (8)

Type : aerobic

Inoculum Contact time

Degradation : (\pm) % after

Result : other: not readily biodegradable

Deg. product

Method : other: calculated using BIOWIN version 4.02

Year : 2006 GLP : no

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Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Remark : Calculation of biodegradation and the timeframe for Primary and Ultimate

biodegradation using BIOWIN version 4.02, a subroutine of the computer program EPI Suite version 3.12 as described by Howard, et al, 1994.

BIOWIN contains six models (linear regression (BIOWIN 1), non-linear regression (BIOWIN 2), ultimate degradation to CO2 and H2O (BIOWIN 3),

primary degradation (BIOWIN 4), linear regression estimate of the probability of passing the OECD 301C / MITI-1 ready biodegradation test (BIOWIN 5), and non-linear regression estimate of the probability of passing the OECD 301C / MITI-1 ready biodegradation test (BIOWIN 6).

BIOWIN 1 - "Does Not Biodegrade Fast" BIOWIN 2 - "Does Not Biodegrade Fast"

BIOWIN 3 - "Weeks" BIOWIN 4 - "Days-Weeks"

BIOWIN 5 - "Does Not Biodegrade Fast" BIOWIN 6 - "Does Not Biodegrade Fast"

According to the USEPA, BIOWIN 6 is better predictor of whether a chemical will pass or fail the OECD 301C / MITI-1 ready biodegradation

test than BIOWIN 5. CAS No. 6863-58-7; sec-butyl ether

Test substance

Reliability : (2) valid with restrictions

The value was calculated based on the chemical structure as modeled by EPI Suite. This robust summary has a reliability rating of 2 because the

data are modeled.

Flag : Critical study for SIDS endpoint

14.11.2006 (6) (7)

Type : aerobic

Inoculum : activated sludge

Concentration: 100 mg/l related to Test substance

related to

Contact time :

Degradation : = 3 - 4 (±) % after 28 day(s) **Result** : other: not readily biodegradable

Deg. product

Method : other: Japanese Guideline by MITI (1974). Comparable to OECD TG 301C,

Modified MITI Test 1

Year : 1992 GLP : no data

Test substance : other TS: CAS No. 6863-58-7; sec-butyl ether

Method : The test was conducted in accordance with "Biodegradation test of

chemical substances by microorganisms etc". stipulated in the Order Prescribing the Items of the Test Relating to the New Chemical Substance (1974, Order of the Prime Minister, the Minister of Health and Welfare, the Minister of International Trade and Industry No 1). This guideline

coresponds to "301C, Ready Biodegradability: Modified MITI Test 1" stipulated in the OECD Guidelines for Testing of Chemicals (1981).

Test condition : Sludge concentration = 30 mg/l

Test substance: Analogue substance CAS No. 142-96-1; n-butyl ether

Reliability : (2) valid with restrictions

Although a standard method was not followed, the testing procedures followed generally accepted aerobic biodegradation guideline methods and although there was limited information on the specifics of the study, there was sufficient information on testing method and conditions in general to

rate this study a 2.

14.11.2006 (2)

ld 6863-58-7 **Date** 14.11.2006

3.6 BOD5, COD OR BOD5/COD RATIO

Species: Cyprinus carpio (Fish, fresh water)

Exposure period : 42 day(s) at °C

Concentration

BCF : = 47 - 83

Elimination

Method : other: guideline corresponds to OECD 305C, Degree of Bioconcentration in

Fish (1981)

Year

GLP : no data

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Result: At a concentration of 0.2 mg/L sec-butyl ether was shown to have

bioconcentrate factor (BCF) range of 47 to 83, which is a log BCF range of

1.67 to 1.92.

At a concentration of 0.02 mg/L sec-butyl ether was shown to have a BCF

range of 30 to 114, which is a log BCF range of 1.48 to 2.06.

Test condition: Fish supplier was Sugishima fish farm, Kumamoto, Japan. Upon arrival,

fish were placed in an acclimation tank with a flow through water system for 28 days at 25° +/-2° C. Test fish were then transferred to test tanks and reared again at the same temperature for approximately 1 month.

At test initiation, average fish weight, length, and lipid content were, 30 g, 10 cm, and 2-5%, respectively. Fish were fed pelleted feed for carp supplied by Haigo Shiryo K.K..

Fish were fed approximately 2% of their total body weight daily. On days fish were sampled, they were not fed.

The test systems included 100 L volume glass tanks with a flow ate of 200 to 800 ml/min at 25° +/-2° C. Dissolved oxygen was 6 to 8 mg/L. At test initiation there was a minimum of 15 fish per exposure level. Treatment levels were based on results of acute toxicity testing.

Test water, test fish, and control fish were analyzed twice a week, every two weeks, and prior to test initiation as well as at termination, respectively. Recovery efficiency was determined in water and fish homogenate. Analytical results from the definitive studies were corrected based on efficiency results.

Bioconcentration factors were calculated based on:

test substance concentration in fish / test substance concentration in water.

Two concentrations were evaluated: 0.2 mg/L and 0.02 mg/L. Analogue substance CAS No. 142-96-1; n-butyl ether

Reliability : (2) valid with restrictions

Although a standard method was not followed, the testing procedures followed generally accepted fish bioconcentration guideline methods and although there was limited information on the specifics of the study, there

was sufficient information on testing method and conditions in general to

rate this study a 2.

Flag : Critical study for SIDS endpoint

14.11.2006 (2)

BCF : = 32.1

Elimination :

Test substance

ld 6863-58-7 **Date** 14.11.2006

Method : other: calculated

Year : 2006 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Method : Calculated values using BCFWIN version 2.15, a subroutine of the

computer program EPIWIN version 3.12.

Test condition :

Test substance

BCFWIN estimates the bioconcentration factor (BCF) of an organic compound using the compound's log octanol-water partition coefficient

(Kow).

The estimation methodology used by BCFWIN is described in "Improved Method for Estimating Bioconcentration Factor (BCF) from Octanol-Water Partition Coefficient", SRC TR-97-006 (2nd Update), July 22, 1997.

Log Kow used = 2.87

Estimated BCF = 32.1 Estimated Log BCF = 1.507

: CAS No. 6863-58-7; sec-butyl ether

Reliability : (2) valid with restrictions

The result is a calculated value based on the chemical structure and represents a potential bioaccumulation factor for the substance with the

CAS number listed under test substance.

14.11.2006 (6)

Type : other: calculated
Species : other: freshwater fish

Exposure period : 96 hour(s)
Unit : mg/l

LC50 : = 14.7 calculated

Method : other: ECOSAR Computer Model

Year : 2006

GLP :

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition: Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA.

Test substance : CAS No. 6863-58-7; sec-butyl ether

Conclusion : Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to

have an acute 96-hour LC50 of 14.7 mg/L and a Chronic Value of 2.2 mg/L

Reliability : (2) valid with restrictions

The value was calculated based on chemical structure as modeled by

EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

Flag : Critical study for SIDS endpoint

14.11.2006 (1)

Type : other: calculated Species : other: freshwater fish

Exposure period : 96 hour(s)
Unit : mg/l

LC50 : = 10.8 calculated

Method : other: ECOSAR Computer Model

Year : 2006 GLP : no

Test substance

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition: Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA. Analogue substance CAS No. 142-96-1; n-butyl ether

Conclusion: Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to have an acute 96-hour LC50 of 10.8 mg/L and a Chronic Value of 1.6

mg/L.

Reliability : (2) valid with restrictions

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

14.11.2006 (1)

Type : semistatic

Species: Oryzias latipes (Fish, fresh water)

Exposure period : 48 hour(s)
Unit : ma/l

LC50 : = 30.7 measured/nominal

Limit test : no Analytical monitoring :

Method: other: Japanese Industrial Standard (JIS K 0102-1986-71): Testing

Methods for Industrial Wastewater.

Year : 1992 GLP : no data

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition: Fish supplier was Nakashima fish farm, Kumamoto, Japan. Upon arrival,

fish were placed in an acclimation tank with a flow-through water system for 3 to 5 weeks after external disinfection. The external disinfection was carried out under static conditions for about 24-hours in an acqueous solution containing 20 mg/l of Elbarju (Ueno Pharm. Co.) and 7 g/l sodium

chloride.

Test fish were then placed in an acclimation tank with a flow-through

system at 25°±2°C for about 28 days.

Dilution water for the test and culture was provided by a well of the Kurume Research Laboratories. Water temperature, pH, and dissolved oxygen were continuously monitored in the laboratory. Total hardness, evaporated residue, chemical oxygen demand, chloride ion, and other harmful substances were also monitored. The quality of the dilution water was confirmed to meet the standards of the Ministry of Health and Welfare in total hardness and evaporated residue. The other analyzed items met the water quality standards for fisheries.

Test tanks were round glass vessels with 4 liters of test water per level. The test was conducted at 25°±2°C. Ten fish were tested at each level for 48 hours under semi-static conditions (renewal of test water every 8 to 16 hours).

Test concentrations (levels) were not reported.

The 48-hour LC50 value was estimated by either the Doudoroff Method or

the Probit Method.

Test substance Reliability

Analogue substance CAS No. 142-96-1; n-butyl ether

(2) valid with restrictions

Although a standard method was not followed, the testing procedures followed generally accepted fish acute toxicity guideline methods and although there was limited information on the specifics of the study, there was sufficient information on testing method and conditions in general to

rate this study a 2.

14.11.2006 (2)

Type :

Id 6863-58-7 4. Ecotoxicity Date 14.11.2006

Species Daphnia sp. (Crustacea)

Exposure period 48 hour(s) Unit mg/l

EC50 = 16.7 calculated

Method other: ECOSAR Computer Model

Year 2006 **GLP**

Test substance other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition : Log Kow (octanol/water partition coefficient) values and a chemical

> structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA.

Test substance Conclusion

Conclusion

CAS No. 6863-58-7; sec-butyl ether

Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to

have an acute 48-hour EC50 of 16.7 mg/L and a Chronic Value of 1.3

mg/L.

(2) valid with restrictions Reliability

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

Critical study for SIDS endpoint Flag

14.11.2006 (1)

Type

Species Daphnia sp. (Crustacea)

Exposure period 48 hour(s) Unit mg/l

EC50 = 12.5 calculated

other: ECOSAR Computer Model Method

Year **GLP** nο

Test substance other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition : Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard, 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA.

Analogue substance CAS No. 142-96-1; n-butyl ether **Test substance**

Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to have an acute 48-hour EC50 of 12.5 mg/L and a Chronic Value of 1.0

mg/L.

(2) valid with restrictions Reliability

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

ld 6863-58-7 4. Ecotoxicity Date 14.11.2006

14.11.2006 (1)

other algae: green alga **Species**

Endpoint

Exposure period 96 hour(s) Unit mg/l

EC50 = 11 calculated

other: ECOSAR Computer Model Method

Year 2006 **GLP** no

Test substance other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition Log Kow (octanol/water partition coefficient) values and a chemical

> structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA.

Test substance : CAS No. 6863-58-7; sec-butyl ether

Conclusion Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to

have an acute 96-hour EC50 of 11.0 mg/L and a Chronic Value of 1.8

mg/L.

Reliability (2) valid with restrictions

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

: Critical study for SIDS endpoint Flag

14.11.2006 (1)

other algae: green alga **Species**

Endpoint

Exposure period 96 hour(s) Unit mg/l

EC50 = 8.3 calculated

Method other: ECOSAR Computer Model

Year 2006 **GLP**

Test substance other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard, 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA.

Test substance Analogue substance CAS No. 142-96-1; n-butyl ether

Conclusion: Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to

have an acute 96-hour EC50 of 8.3 mg/L and a Chronic Value of 1.5 mg/L.

Reliability : (2) valid with restrictions

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

14.11.2006 (1)

Species: other: Freshwater Fish (calculated toxicity values are not species specific)

Endpoint : other: LC50
Exposure period : 30 day(s)
Unit : mg/l

ChV : = 2.2 calculated

Method : other: ECOSAR Computer Model

Year : 2006 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition: Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA.

Test substance: CAS No. 6863-58-7; sec-butyl ether

Conclusion: Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to

have an acute 96-hour LC50 of 14.7 mg/L and a Chronic Value of 2.2 mg/L

Reliability : (2) valid with restrictions

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

Flag : Critical study for SIDS endpoint

14.11.2006 (1)

Species: other: Freshwater Fish (calculated toxicity values are not species specific)

Endpoint : other: LC50
Exposure period : 30 day(s)
Unit : mg/l

ChV : = 1.6 calculated

Method : other: ECOSAR Computer Model

Year : 2006 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition : Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

Test substance : Analogue substance CAS No. 142-96-1; n-butyl ether

Conclusion : Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to

have an acute 96-hour LC50 of 10.8 mg/L and a Chronic Value of 1.6

mg/L.

Reliability : (2) valid with restrictions

The value was calculated based on chemical structure as modeled by

EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

14.11.2006 (1)

Species : Daphnia sp. (Crustacea)

Endpoint : mortality
Exposure period : 16 day(s)
Unit : mg/l

EC50 : = 1.3 calculated

Method : other: ECOSAR Computer model

Year : 2006 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition: Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.

Syracuse Research Corporation, Syracuse, NY, USA.

Test substance: CAS No. 6863-58-7; sec-butyl ether

Conclusion: Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to

have an acute 48-hour EC50 of 16.7 mg/L and a Chronic Value of 1.3

mg/L.

Reliability : (2) valid with restrictions

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

Flag : Critical study for SIDS endpoint

14.11.2006 (1)

Species : Daphnia sp. (Crustacea)

Endpoint : mortality
Exposure period : 16 day(s)
Unit : mg/l

EC50 : = 1 - calculated

Method : other: ECOSAR Computer model

Year : 2006 GLP : no

Test substance: other TS: CAS No. 6863-58-7; sec-butyl ether

Test condition : Log Kow (octanol/water partition coefficient) values and a chemical

structure are needed to calculate aquatic toxicity using the ECOSAR model. The Kow calculation is performed by KOWWIN based on an atom/fragment contribution method of Meylan and Howard (1), which is a

subroutine in the EPISuite computer model (2).

1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.

2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12. Syracuse Research Corporation, Syracuse, NY, USA.

Analogue substance CAS No. 142-96-1; n-butyl ether

: Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to have an acute 48-hour EC50 of 12.5 mg/L and a Chronic Value of 1.0

mg/L.

Reliability : (2) valid with restrictions

Test substance

Conclusion

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

14.11.2006 (1)

5. Toxicity	ld	ld 6863-58-7	
	Date	14.11.2006	
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5. Toxicity	D	6863-58-7 14.11.2006
5.11 ADDITIONAL REMARKS		
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6. Analyt. Meth. for Detection and Identification	6863-58-7 14.11.2006
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7. Eff. Against Target Org. and Intended Uses	6863-58-7 14.11.2006
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9. References Id 6863-58-7 Date 14.11.2006

(1) Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment Division. Washington, DC, USA.

- (2) Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation and bioaccumulation data of existing chemicals based on the CSCL Japan. CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau, Ministry of International Trade and Industry, Japan. Japan Chemical Industry Ecology-Toxicology and Information Center.
- (3) Drake, Nathan L. (1935). Journal of the American Chemical Society. Vol 57, pp. 2623-2625 CAPLUS.
- (4) EMBSI (2006) Hydrolysis: CAS No. 6863-58-7; sec-Butyl Ether.
- (5) EMBSI (2006) Photodegradation (Direct): CAS No. 6863-58-7; sec-Butyl Ether.
- (6) EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12. Syracuse Research Corporation, Syracuse, NY, USA.
- (7) Howard, PH, Boethling, RS, Stitler, WM, Meylan, WM, Hueber, AE, Beauman, JA and Larosche, ME (1992). Predictive model for aerobic biodegradability developed from a file of evaluated biodegradation data. Environ Toxicol Chem 11, 593-603.
- (8) Mackay D, DiGuardo A, Paterson S and Cowan C (2003). EQC Model ver. 2.02, available from the Environmental Centre, Trent University, Canada.

10. Summary and Evaluation	6863-58-7 14.11.2006
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2007 JAN -3 AM 7: 31

201-16472C

IUCLID

Data Set

Existing Chemical

CAS No.

; ID: 108-20-3 : 108-20-3

EINECS Name

: diisopropyl ether

EC No.

: 203-560-6

: Propane, 2,2'-oxybis-

TSCA Name Molecular Formula

: C6H14O

Producer related part

Company

: ExxonMobil Biomedical Sciences Inc.

Creation date

: 18.05.2005

Substance related part

Company

: ExxonMobil Biomedical Sciences Inc.

Creation date

: 18.05.2005

Status

Memo

: HPV

Printing date

: 27.02.2006

Revision date

Date of last update

: 27.02.2006

Number of pages

: 55

Chapter (profile) Reliability (profile) : Chapter: 1, 2, 3, 4, 5, 6, 7, 8, 10

Flags (profile)

: Reliability: without reliability, 1, 2, 3, 4

: Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE),

Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

1. General Information

Id 108-20-3

Date

1.0.1 APPLICANT AND COMPANY INFORMATION

1.0.2 LOCATION OF PRODUCTION SITE, IMPORTER OR FORMULATOR

1.0.3 IDENTITY OF RECIPIENTS

1.0.4 DETAILS ON CATEGORY/TEMPLATE

1.1.0 SUBSTANCE IDENTIFICATION

1.1.1 GENERAL SUBSTANCE INFORMATION

Purity type :

Substance type : organic Physical status : liquid

Purity : Colour : Odour :

27.10.2005

1.1.2 SPECTRA

1.2 SYNONYMS AND TRADENAMES

2,2'-oxybis-propane

27.10.2005

2,2'-oxybispropane

27.10.2005

2-Isopropoxy Propane

27.10.2005

2-Isopropoxypropan

27.10.2005

2-isopropoxypropane

27.10.2005

Diisopropyl Ether

27.10.2005

Id 108-20-3 1. General Information **Date** Dipropyloxid 27.10.2005 IPE 27.10.2005 IPE; Diisopropylether; DIPE; 2-Isopropoxy propane 27.10.2005 **Isopropyl Ether** 27.10.2005 Isopropylether 27.10.2005 propane, 2,2'-oxybis-27.10.2005 1.3 IMPURITIES 1.4 ADDITIVES 1.5 TOTAL QUANTITY 1.6.1 LABELLING 1.6.2 CLASSIFICATION 1.6.3 PACKAGING 1.7 USE PATTERN 1.7.1 DETAILED USE PATTERN 1.7.2 METHODS OF MANUFACTURE 1.8 REGULATORY MEASURES

Date 27.02.2006 1.8.1 OCCUPATIONAL EXPOSURE LIMIT VALUES 1.8.2 ACCEPTABLE RESIDUES LEVELS 1.8.3 WATER POLLUTION 1.8.4 MAJOR ACCIDENT HAZARDS 1.8.5 AIR POLLUTION 1.8.6 LISTINGS E.G. CHEMICAL INVENTORIES 1.9.1 DEGRADATION/TRANSFORMATION PRODUCTS 1.9.2 COMPONENTS 1.10 SOURCE OF EXPOSURE 1.11 ADDITIONAL REMARKS 1.12 LAST LITERATURE SEARCH 1.13 REVIEWS

1. General Information

Id 108-20-3

ld 108-20-3

Date

2.1 MELTING POINT

Value : = -86.8 °C

Sublimation

Method : other: not specified

Year

GLP : no data

Test substance : other TS: Diisopropyl Ether (CAS # 108-20-3)

Test substance : CAS No. 108-20-3; diisopropyl ether; purity is unknown.

Reliability : (2) valid with restrictions

The CRC Handbook of Chemistry and Physics is a peer reviewed

publication. This robust summary has a reliability rating of 2 because there is insufficient information available on the method and analytical procedure.

Flag : Critical study for SIDS endpoint

07.12.2005 (25)

2.2 BOILING POINT

Value : = 68.5 °C at 1013 hPa

Decomposition

Method : other: not specified

Year

GLP : no data

Test substance : other TS: Diisopropylether

Test substance: CAS No. 108-20-3; diisopropyl ether; purity is unknown.

Reliability : (2) valid with restrictions

The CRC Handbook of Chemistry and Physics is a peer reviewed

publication. This robust summary has a reliability rating of 2 because there is insufficient information available on the method and analytical procedure.

Flag : Critical study for SIDS endpoint

27.10.2005 (25)

2.3 DENSITY

Type : density

Value : = .7241 g/cm³ at 20 °C Method : other: not specified

Year

GLP : no data

Test substance : other TS: Diisopropyl Ether (CAS # 108-20-3)

Test substance: CAS No. 108-20-3; diisopropyl ether; purity is unknown.

Reliability : (2) valid with restrictions

The CRC Handbook of Chemistry and Physics is a peer reviewed

publication. This robust summary has a reliability rating of 2 because there is insufficient information available on the method and analytical procedure.

Flag : Critical study for SIDS endpoint

07.12.2005 (25)

2.3.1 GRANULOMETRY

2. Physico-Chemical Data

ld 108-20-3

Date

2.4 VAPOUR PRESSURE

Value : = 198.65 hPa at 25 °C

Decomposition Method Year

GLP : no data

Test substance: other TS: Diisopropyl Ether (CAS # 108-20-3)

Method: Method not specified.

Test substance : CAS No. 108-20-3; diisopropyl ether; purity is unknown.

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data were not

reviewed for quality, however, the reference is from a peer-reviewed

handbook

Flag : Critical study for SIDS endpoint

07.12.2005 (9)

2.5 PARTITION COEFFICIENT

Partition coefficient : octanol-water Log pow : = 1.52 at 25 °C

pH value

Method : other (measured)

Year

GLP : no data

Test substance : other TS: Diisopropylether

Method: Method not specified.

Test substance: CAS No. 108-20-3; diisopropyl ether; purity is unknown.

Reliability : (2) valid with restrictions

The value cited by the authors is a measured and preferred value. This robust summary has a reliability rating of 2 because there is insufficient

information available on the method and analytical procedure.

Flag : Critical study for SIDS endpoint

27.10.2005 (19)

Partition coefficient : octanol-water Log pow : = 2.4 at °C

pH value : 6.7

Method : other (calculated): Indirect method by reverse-phase HPLC

Year :

GLP : no

Test substance : other TS: diisopropyl ether (CAS No. 108-20-3)

Result : Log Pow = 2.4 (Pow = 250) at pH 6.7

Test condition: The HPLC system was a reverse-phase C18-coated silica gel column, 250

mm x 5 mm id, with a mobile phase of 3 volumes methanol and 1 volume water (final pH 6.7) at a flow rate of 1 mL/min. 100 mL samples of an approximate 1 mg/mL solution in the mobile phase were injected, and the emergence of the material was observed using refraction index detection. Thirty-one reference compounds were used to generate the linear relationship between log k (k = capacity factor) and log Pow. Using the HPLC retention time for the peak of the test substance, the log k was

determined, and the log Pow value was calculated using the linear equation developed from the reference compounds.

Log Pow was determined according to the following calculations:

Retention time (RT), min = 5.7

2. Physico-Chemical Data

ld 108-20-3

Date

Capacity factor, k = 0.87, k = (RTcmpd - RTunretained std)/RTunretained

std

 $\log k = -0.06$

linear equation: $\log k = -0.930 + 0.357 \log Pow$

Reliability : (1) valid without restriction

12.12.2005 (11)

2.6.1 SOLUBILITY IN DIFFERENT MEDIA

Solubility in : Water

Value : = 8800 mg/l at 20 °C

pH value

concentration : at °C

Temperature effects

Examine different pol.

pKa : at 25 °C

Description :
Stable :
Deg. product :
Method :

Year

GLP : no data

Test substance : other TS: Diisopropyl Ether (CAS # 108-20-3)

Test substance: CAS No. 108-20-3; diisopropyl ether; purity is unknown.

Reliability : (2) valid with restrictions

The Ullmann's Encyclopedia of Industrial Chemistry is a peer reviewed publication. This robust summary has a reliability rating of 2 because there is insufficient information available on the method and analytical procedure.

Flag : Critical study for SIDS endpoint

07.12.2005 (17)

2.6.2 SURFACE TENSION

2.7 FLASH POINT

2.8 AUTO FLAMMABILITY

2.9 FLAMMABILITY

2.10 EXPLOSIVE PROPERTIES

2.11 OXIDIZING PROPERTIES

2.12 DISSOCIATION CONSTANT

2.13 VISCOSITY

2. Physico-Chemical Data	Id Date	108-20-3
2.14 ADDITIONAL REMARKS		
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ld 108-20-3

Date

3.1.1 PHOTODEGRADATION

Type : air Light source :

Light spectrum : nm

Relative intensity: based on intensity of sunlight

Conc. of substance : at 25 °C

INDIRECT PHOTOLYSIS

Sensitizer : OH

Conc. of sensitizer : 1500000 molecule/cm³

Rate constant : $= .00000000002434 \text{ cm}^3/(\text{molecule*sec})$

Degradation : = 50 % after 5.3 hour(s)

Deg. product

Method : other (calculated): Calculated values using AOPWIN version 1.89, a

subroutine of the computer program EPIWIN version 3.12

Year

GLP

Test substance: other TS: Diisopropyl Ether (CAS # 108-20-3)

Method : Calculated values using AOPWIN version 1.89, a subroutine of the

computer program EPIWIN version 3.12

Indirect photodegradation, or atmospheric oxidation potential, is based on the structure-activity relationship methods developed by R. Atkinson under

the following conditions: Temperature: 25°C Sensitizer: OH- radical

Concentration of Sensitizer: 1.5E6 OH- radicals/cm3

Remark : DIPE has the potential to volatilize to air, based on a vapor pressure of

19,865 Pa at 25°C (Daubert and Danner, 1989), where it is subject to atmospheric oxidation. In air, DIPE can react with photosensitized oxygen in the form of hydroxyl radicals (OH-). The computer program AOPWIN (atmospheric oxidation program for Microsoft Windows) (EPIWIN, 2000) calculates a chemical half-life for a 12-hour day (the 12-hour day half-life value normalizes degradation to a standard day light period during which hydroxyl radicals needed for degradation are generated), based on an OH-

reaction rate constant and a defined OH- concentration.

DIPE has a calculated half-life in air of 5.3 hours or 0.4 days (12-hour day), based on a rate constant of 24.34 E-12 cm3/molecule*sec and an OH-

concentration of 1.5 E5 OH-/cm3.

Reliability : (2) valid with restrictions

The value was calculated based on chemical structure as modeled by EPIWIN. This robust summary has a reliability rating of 2 because the data

are calculated and not measured.

Flag : Critical study for SIDS endpoint

07.12.2005 (15)

Deg. product : Method : Year : GLP :

Test substance : other TS: Diisopropyl Ether (CAS # 108-20-3)

Method : Technical discussion

Remark: Direct photochemical degradation occurs through the absorbance of solar

radiation by a chemical substance in aqueous solution. If the absorbed energy is high enough, then the resultant excited state of the chemical may undergo a transformation. A prerequisite for direct photodegradation is the

ability of one or more bonds within a chemical to absorb ultraviolet

Id 108-20-3

Date

(UV)/visible light in the 290 to 750 nm range. Light wavelengths longer than 750 nm do not contain sufficient energy to break chemical bonds, and wavelengths below 290 nm are shielded from the earth by the stratospheric ozone layer (Harris, 1982a).

An approach to assessing the potential for a substance to undergo photochemical degradation is to assume that degradation will occur in proportion to the amount of light wavelengths >290 nm absorbed by constituent molecules (Zepp and Cline, 1977). The oxygen non-bonding electrons in ethers do not give rise to absorption above 160 nm, which is

why pure ether solvents can be used in spectroscopic studies.

Consequently, DIPE is not subject to photolytic processes in the aqueous

environment.

Reliability (2) valid with restrictions

This robust summary has a reliability of 2 because it is a technical

discussion and not a study.

: Critical study for SIDS endpoint Flag

07.12.2005 (46)

3.1.2 STABILITY IN WATER

Type abiotic at °C t1/2 pH4 at °C t1/2 pH7 at °C t1/2 pH9

Deg. product

other: Technical discussion Method

Year

GLP no data

Test substance other TS: Diisopropyl Ether (CAS # 108-20-3)

Result Hydrolysis of an organic chemical is the transformation process in which a

water molecule or hydroxide ion reacts to form a new carbon-oxygen bond. Chemicals with leaving groups that have a potential to hydrolyze include alkyl halides, amides, carbamates, carboxylic acid esters and lactones, epoxides, phosphate esters, and sulfonic acid esters (Gould, 1959). The lack of a suitable leaving group renders a compound resistant to hydrolysis. DIPE is resistant to hydrolysis because it lacks a functional group that is hydrolytically reactive and Harris (1982b) identifies ether groups as

generally resistant to hydrolysis. Therefore, hydrolysis will not contribute to

the removal of diisopropyl ether from the environment.

: (2) valid with restrictions Reliability

This robust summary has a reliability of 2 because it is a technical

discussion and not a study.

: Critical study for SIDS endpoint Flag

07.12.2005 (18)(20)

3.1.3 STABILITY IN SOIL

3.2.1 MONITORING DATA

3.2.2 FIELD STUDIES

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

ld 108-20-3

Date

Туре

Media : other: air - biota - sediment(s) - soil - water

Air : % (Fugacity Model Level I)
Water : % (Fugacity Model Level I)
Soil : % (Fugacity Model Level I)
Biota : % (Fugacity Model Level II/III)
Soil : % (Fugacity Model Level II/III)

Method : other: Calculation according Mackay, Level I

Year

Remark: Physicochemical data used in the calculation:

Parameter Value w/ Units

Molecular Weight = 102.18 Temperature = 25° C Log Kow = 1.52

Water Solubility = 8,800 g/m3 Vapor Pressure = 19,865 Pa Melting Point = -86.8° C

Result : Using the Mackay Level I calculation, the following

distribution is predicted for diisopropyl ether:

%Distribution Compartment

97.83 Air 2.10 Water 0.06 Soil <0.01 Sediment

< 0.01 Suspended Sediment

<0.01 Biota

Test substance : Diisopropyl Ether (CAS # 108-20-3)

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated.

Flag : Critical study for SIDS endpoint

07.12.2005

Type : fugacity model level III

Media : other

Air : % (Fugacity Model Level I)

Water : % (Fugacity Model Level I)

Soil : % (Fugacity Model Level I)

Biota : % (Fugacity Model Level II/III)

Soil : % (Fugacity Model Level II/III)

Method : other: Level III simulation using the Mackay Multimedia Environmental

Model (Mackay, 2001)

Year :

Method : Level III simulation using the Mackay Multimedia Environmental Model

(Mackay, 2001). Mass balances are calculated for the four bulk media of air (gas + aerosol), water (solution + suspended sediment + biota), soil, (solids + air + water), and sediment (solids + pore water). Equilibrium exists within, but not between media. Physical-chemical properties are used to quantify a

chemical's behavior in an evaluative environment. Three types of

chemicals are treated in this model: chemicals that partition into all media (Type 1), non volatile chemicals (Type 2), and chemicals with zero, or near-zero, solubility (Type 3). The model can not treat ionizing or speciating substances. The Level III model assumes a simple, evaluative environment with user-defined volumes and densities for the following homogeneous environmental media (or compartments): air, water, soil, sediment,

suspended sediment, fish and aerosols.

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This model provides a description of a chemical's fate including the important degradation and advection losses and the intermedia transport processes. The distribution of the chemical between media depends on how the chemical enters the system, e.g. to air, to water, or to both. This mode of entry also affects persistence or residence time.

The rates of intermedia transport are controlled by a series of 12 transport velocities. Reaction half-lives are requested for all 7 media. The advective residence time selected for air also applies to aerosols and the residence time for water applies to suspended sediment and fish. The advective residence time of aerosols, suspended sediment and fish cannot be specified independently of the air and water residence times.

Result : Output

Output			
-	Mass%	Half life(hr)	Emissions(kg/hr)
Air	19.4	25.2	1000
Water	61.0	360	1000
Soil	19.5	720	1000
Sediment	0.1	3240	0

Test condition : Physchem Inputs

Molar Mass = 102.18 Data Temperature = 25 °C Water Solubility = 8800 mg/l exp. Vapour Pressure = 19865 Pa exp.

Log Kow = 1.52 exp.

Melting Point = -86.8 °C exp.

Reaction Half Lives in hours (if not available they can be predicted using EPIWIN)

Air (gaseous) 25.2
Water (no susp. part.) 360
Bulk Soil 720
Bulk Sediment 3240
Suspended Particles 360
Fish 360
Aerosol 25.2

Environmental Properties (EQC standard environment)

Dimensions (all defaults) Densities (all defaults)

Organic carbon & Advection (all defaults)

Transport Velocities (all defaults)

Emission and Inflows (defaults used)

Air 1000 kg/hr Water 1000 kg/hr Soil 1000 kg/hr Sediment 0 kg/hr

Test substance: Diisopropyl Ether, CAS No. 108-20-3

: The majority of DIPE is calculated to partition into the water phase, with smaller but significant amounts into air and soil, based on the modeling parameters used in this calculation. DIPE is considered to be a Type 1

chemical with potential to partition into all environmental compartments.

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated.

Flag : Critical study for SIDS endpoint

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3.3.2 DISTRIBUTION

Conclusion

Id 108-20-3

Date

3.4 MODE OF DEGRADATION IN ACTUAL USE

3.5 BIODEGRADATION

Type : aerobic

Inoculum : activated sludge, domestic

Contact time : 28 day(s)
Degradation : (±) % after

Result : other: not readily biodegradable

Deg. product

Method : OECD Guide-line 301 D "Ready Biodegradability: Closed Bottle Test"

Year : 1982 **GLP** : no

Test substance: other TS: Diisopropyl Ether (CAS No. 108-20-3)

Result: Test substance was not readily biodegradable. After 28 days, the test

substance exhibited no measurable biodegradation. By day 5, >60% biodegradation of positive control was observed, which meets the guideline requirement. No excursions from the testing guideline were noted. The inhibition study showed that the test substance did not inhibit the biodegradability of the positive control substance, sodium benzoate.

Sample	% Degradation* Mean (day 28)	% Degradation (day 28)
Test Substance Na Benzoate * duplicate data	65.0, 73.0	0.0 69.0

Mean oxygen concentrations (mg/L) of duplicate test systems:

Day 0

Mineral Salts Control = 8.85

Blank = 8.8

Na Benzoate = 8.95

Test Substance = 8.9 (single test system)

Test Substance + Na Benzoate = 8.9* (single test system)

Day 5

Mineral Salts Control = 9.0

Blank = 8.8 Na Benzoate = 5.7 Test Substance = 8..85

Test Substance + Na Benzoate = 5.8

Day 15

Mineral Salts Control = 8.75

Blank = 8.65 Na Benzoate = 4.9 Test Substance = 8.55

Test Substance + Na Benzoate = 4.9

Day 28

Mineral Salts Control = 8.65

Blank = 7.05 Na Benzoate = 3.6 Test Substance = 8.3

Test Substance + Na Benzoate = 4.15

Test condition: The inoculum source was the Sittingbourne Sewage works in Kent,

ld 108-20-3

Date

England, and was prepared according to methods described in the OECD 301D guideline. The test substance was added to the test medium by direct addition at a concentration of 3.0 mg/L. Test systems were incubated at 20 \pm 1 °C and biodegradation was determined by measuring the oxygen concentration on days 5, 15, and 28. Each sampling of the test substance and control was conducted in duplicate. The theoretical oxygen demand was 2.82 mg O2 per mg test substance and a theoretical carbon dioxide (CO2) evolution of 2.59 mg CO2 per mg test substance. Sodium benzoate was used as the positive control.

The purity of the test substance was not supplied, but the infra-red spectrum of the test substance matched a published standard (density = 0.723 to 0.726 kg/L). The test substance was stored in the dark at ambient temperature. Nitrogen was blown over the surface of the material when the container was opened and exposed to air in order to minimize peroxide formation.

: Diisopropyl ether is not readily biodegradable and it did not significantly

inhibit the biodegradability of the test substance in an inhibition test.

Reliability : (1) valid without restriction

07.12.2005 (40)

Type

Conclusion

Inoculum : other: sanitary waste treatment plant effluent

Contact time : 5 day(s)
Degradation : (±) % after

Result

Deg. product

Method : other: American Public Health Association; No. 219 5-Day BOD; Standard

Dilution Method

Year : 1971 **GLP** : no

Test substance : other TS: diisopropyl ether (CAS No. 108-20-3)

Remark: Test type: Biological Oxygen Demand (BOD)

Result : $0.19 \text{ g O2/g test material at } 20 \pm 1^{\circ}\text{C}$

The theoretical oxygen demand (ThOD) of the test substance was 2.82.

The percent ThOD in 5 days was 7%.

The article stated that the only deviation from the standard method was the

addition of 0.5 mg/L allylthiourea to prevent nitrification.

Test condition: The article stated that the test method followed APHA Standard Method

No. 219 (1971). The test was run at a temperature of $20 \pm 1^{\circ}$ C. 500-mL test solutions were seeded with a filtered 10-mL volume of the effluent from a biological sanitary waste treatment plant. The activity of the inoculum was check by including a treatment containing a mixture of glucose and glutamic acid. Test mixtures were stirred using a magnetic stirrer.

Conclusion : 5-day BOD = 0.19 g/g, representing 7% biodegradation of the test

substance.

Reliability : (2) valid with restrictions

The article presented a brief description of the testing methods, but cited a

reliable guideline method in use at the time of the study.

07.02.2006 (2)

Type : aerobic

Inoculum : activated sludge

Deg. product

Method: other: (comparison study of three aerobic biodegradation methods)

Year : 1997 **GLP** : no

Test substance: other TS: diisopropyl ether (CAS No. 108-20-3)

Method : Comparison study of three aerobic biodegradation methods)

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Date

Continuous Biological Treatment: (1) EPA Method 304B (EPA, 1994)

Batch Methods:

(2) Batch Oxygen addition (BOX) (Rajagopalan et al., 1998), and

(3) Serum Bottle Test (SBT) (Rajagopalan et al., 1998)

Remark : Exposure Period:

Method 304: 30 days BOX: 0.5 to 2 hours SBT: 0.5 to 2 hours

Result : The average percent removal of the test substance in the continuous

activated sludge unit (EPA Method 304B) was 99.4%.

Three experimental trial runs with each of the three biodegradation methods yielded the following average first-order biodegradation rate constants (K1 = L/g Volatile Suspended Solids-h) for the test substance:

K1 (L/g VSS-h)

304B 98 BOX 17.4 SBT 19.2

Test condition

A pilot-scale continuous activated sludge unit served as the source of biomass for kinetic rate constant comparisons of the three methods. The activated sludge was acclimated in the pilot unit by feeding a synthetic cocktail of eight volatile organic compounds during a 2-month equilibration period. Equilibration and testing was done at ambient temperature (22 to 24°C). The hydraulic retention time (HRT) was 7.7 hours and the solids retention time (SRT) was 33 days. Average organic removal efficiencies based on COD and TOC were 92 and 88%, respectively.

During the biodegradation testing using Method 304B, feed and effluent samples were collected in headspace-free VOA vials and stored at 4°C until analyzed. Samples were analyzed by purge-and-trap gas chromatography using a flame ionization detector. Triplicate biodegradation runs on the test compound were conducted with at least six influent and effluent samples taken at 1 HRT (approx. 8 hours) intervals.

The two batch biodegradation testing methods (BOX and SBT) used activated sludge biomass from the pilot-scale reactor. Biomass was diluted using effluent from the system to achieve range of 200 to 600 mg/L. The test compound was injected into the batch reactors and the concentration was monitored over time by collecting gas samples directly from the headspace using an automatic sampling pump and analyzing immediately using gas chromatography.

Conclusion

The authors indicated that K1 values >10 L/g VSS-h represent readily biodegradable organic compounds. Based on the results of this study, all three test methodologies showed the test substance to be effectively utilized by activated sludge microorganisms under aerobic conditions.

Reliability : (2) valid with restrictions

The publication presented a well-documented study based on ound

scientific principles.

07.02.2006 (6) (34) (42)

Type : aerobic

inoculum : other: Mixture (see remarks)

Contact time : 600 day(s)

Degradation : (±) % after

Result Deg. product

Method : other: (continuous-flow bioreactors)

Year : 2001 GLP : no

Test substance : other TS: diisopropyl ether (CAS No. 108-20-3)

Result

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Remark: Inoculum consisted of a mixture of the following:

1) mixed liquor from the Metropolitan Sewer District (MSD), Cincinnati, OH,

2) mixed liquor from Shell Development Co., Houston, TX, and

3) aquifer material wash water from a MTBE-contaminated site in Port

Hueneme, CA.

: The authors indicated that removal of DIPE was comparable to that

achieved for MTBE, which was greater than 99.9%.

Test condition: Bioreactors (Autoclave Engineers, Erie, PA) were initially seeded with 2 L

of mixed liquor from the MSD, 600 mL of mixed liquor from Shell Development Co., and 140 mL of aquifer wash water. Cultures were maintained on a total influent feed of 417 mg/L chemical oxygen demand (COD) divided as 50% methyl tert-butyl ether (MTBE) and 50% as

diisopropyl ether (DIPE).

Reactors were well mixed and controlled to a temperature of 20°C. To retain high biomass levels, a polyethylene porous pot was inserted into the reactor. The pots consisted of 0.45 cm thick filter-grade polyethylene (pore size = 20 mm), with an internal diameter of 19.1 cm and a height of 29.2 cm. Initially, a solids retention time of 18 days was maintained by wasting intentionally from the reactor. Subsequently (after about 120 days) intentional wasting ceased and only took place during sampling of the reactors.

The combined influent flow rate was 2.37 L/d, with 80% of the total flow provided by a pH-adjustment solution, and 20% provided by an acidified nutrient solution. The pH-adjustment solutions contained deionized water, MTBE and DIPE fed by a syringe infusion pump, and an appropriate amount of 10N sodium hydroxide to maintain the pH between 7.4 and 8.0. The nutrient solution consisted of deionized water with essential salts and vitamins added to promote biological growth. Final nutrient concentrations inside the reactor were as follows: (NH4)2SO4, 93 mg/L; MgSO4, 69.6 mg/L; CaCl2o2H2O, 22.5 mg/L; K2HPO4, 6.9 mg/L; CuSO4oH2O, 0.08 mg/L; Na2MoO4o2H2O, 0.15 mg/L; MnSO4oH2O, 0.13 mg/L; ZnCl2, 0.23 mg/L; CoCl2o6H2O, 0.42 mg/L; and FeCl2o4H2O, 17.25 mg/L.. The hydraulic retention time was 4.2 days with a total reactor volume of 9.95 L and an enrichment culture volume of 6 L.

Effluent from the reactors was monitored weekly for the presence of MTBE and DIPE using gas chromatography equipped with a flame ionization detector (FID) and a 60/80 Carbopack B5% Carbowax 20 M glass column. The pH of the system was measured daily, and COD and dissolved organic carbon (DOC) was measured weekly.

: Diisopropyl ether can be effectively biodegraded in high biomass aerobic

reactors.

Reliability : (2) valid with restrictions

The report provided adequate details of the test conditions but reported

only a text description of biodegradation results.

07.02.2006 (33)

Type :

Conclusion

Inoculum : other: soil and groundwater from a site previously exposed to methyl tert-

butyl ether

Contact time : 1 year
Degradation : (±) % after

Result :

Method : other: (soil/water microcosm)

Year : 1999 **GLP** : no

Test substance: other TS: diisopropyl ether (CAS No. 108-20-3)

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Remark : Test type: soil/water microcosm

Result: No detectable biodegradation of the test substance occurred after one year

of incubation.

Test condition : Soil and water from an aquifer with previous exposure to methyl tert-butyl

ether (MTBE) was collected using a coring device and a pump. The material was brought to the laboratory where the sediment was thoroughly mixed. Groundwater was filtered through a 0.45 mm filter and sparged for 12 hours with sterile air to oxygenate the water and to remove background volatile chemicals. Analysis by gas chromatography indicated that concentrations of MTBE in the aqueous samples were <10 mg/L. Microcosms were constructed in amber 255-mL screw-top bottles sealed with TeflonÒ MininertÒ valves. Each bottle contained 150 g of wet sediment, 140 mL of sterile groundwater and 3000 mg/L of diisopropyl ether (DIPE). Treatments were constructed in triplicate with matching abiotic controls. Sediment used for the abiotic controls was autoclaved for one hour on each of three consecutive days. Additionally, 250 mg/L of mercuric chloride was added to ensure no biological activity. Microcosms were incubated in the dark at 16 °C.

All samples were analyzed every 30 days by purge and trap gas chromatography and flame ionization detection to determine concentrations of the test substance. Test substance disappearance relative to abiotic controls was the principal indicator of biodegradation.

Conclusion: The test substance was not aerobically biodegraded by indigenous

subsurface microflora.

Reliability : (2) valid with restrictions

The testing method did not follow any specific regulatory guideline method, but the publication provided valuable information using sound scientific

principles.

07.02.2006 (45)

Type : anaerobic

Inoculum : other: sediment and groundwater from an anoxic aquifer polluted by

municipal landfill leachate

Contact time : 252 day(s)
Degradation : (±) % after

Result : Deg. product :

Method : other: (closed serum bottle test)

Year : 1993 **GLP** : no

Test substance: other TS: diisopropyl ether (CAS No. 108-20-3)

Result : Biodegradation Rate (ppm C/day) = 0

Methane recovery (% theoretical) = 0

Test condition : Diisopropyl ether was tested for the ability of the compound to be

completely biodegraded to methane in an aquifer slurry. Sediment and groundwater were collected from a methanogenic portion of a shallow anoxic aquifer polluted by municipal landfill leachate. Slurries were prepared by placing 50 g of sediment and 75 mL of groundwater in sterile 160-mL serum bottles. The bottles were sealed with Teflon-lined stoppers and incubated in the dark at room temperature. Diisopropyl ether was added to the incubation mixture to reach an initial substrate concentration of 50 ppm C. Pressure increases resulting from biogas formation (CH4 and CO2) were monitored with an automated pressure transducer system. The acclimation time was estimated as the amount of time where no significant pressure difference was measured between the substrate-

amended treatment and un-amended controls.

At the end of the incubation period, biodegradation was measured as the depletion of parent substrate and the formation of methane over background controls. Measurements were made using gas

Date

chromatography equipped with a flame ionization detector. A 1.8 m x 0.32 cm 80/100 porapak Q column or a 0.2% Carbowax 1500 on Carbopack C column were used for headspace methane analyses and test substance determinations, respectively. Autoclaved controls were similarly assayed and were uniformly unable to exhibit methane formation or test substance disappearance. The amount of methane formed in aquifer incubations was compared to that theoretically expected based on the Buswell equation.

Conclusion : Diisopropyl ether was a persistent molecule that resisted anaerobic

destruction. After 252 days, no evidence for the anaerobic biodegradability

of diisopropyl ether was obtained.

Reliability : (2) valid with restrictions

The publication reported a well-documented study that meets basic

scientific principles.

07.02.2006 (41)

Type : anaerobic

Inoculum : other: Sediment and surface or groundwater

Deg. product

Method : other: unknown

Year : 1994 **GLP** : no

Test substance : other TS: diisopropyl ether (CAS No. 108-20-3)

Remark: Exposure period: 85, 180, or 244 days

Result : Biodegradation of diisopropyl ether with sulfate or nitrate available as

electron acceptors:

SO4 or NO3

Substrate Amount Consumed Rate Loss (%) (% Theoretical) (umol/SO4/day)

sulfate-reducing 0 0 0 0 nitrate-reducing 0 0 0

Biodegradation of diisopropyl ether under methanogenic conditions:

Degradation Methane Recovery Rate (ppm C/day) (% Expected)

Fuel-impacted river

sediment 0 0

Industrial/sewage

impacted creek sediment 0 6

Test condition

Several tests were carried out to determine the anaerobic biodegradation of the test substance. Three experiments were done to determine biodegradation under sulfate- and nitrate-reducing conditions and under methanogenic conditions. Sediment and surface water (or groundwater) from three sources were used as inoculum in separate experiments; (1) sediment/groundwater from a landfill leachate impacted aquifer, (2) sediment/surface water from a river historically impacted by oil storage and barge loading facilities, and (3) sediment/surface water from a creek impacted by industrial waste and domestic sewage sludge.

Slurries were prepared by placing 50 g of sediment and 75 mL of water into sterile 160-mL serum bottles. Water was amended with sodium sulfide (1 mM) and resazurin (0.0002%) to serve as reductant and redox indicator, respectively. The bottles were sealed with stoppers and the headspace above the slurries was adjusted to 80% N2:20%CO2 (1 atm). To the landfill leachate-impacted samples, either sodium sulfate (5mM) or sodium nitrate (8 mM) was added in order to assess potential test substance decay coupled with the consumption of these electrons (referred to as sulfate-reducing and nitrate-reducing incubations, respectively). The test substance was added to the slurries to give an initial concentration of 50 ppm C. The rates of methane production, sulfate reduction, and nitrate

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depletion were monitored in slurries receiving the test substance and compared to test substance-free controls. All incubations were done in the dark at 24°C. The sulfate-reducing experiment was run for 244 days, the nitrate-reducing experiment was run for 85 days, and the methanogenic experiment was run for 180 days.

In the methanogenic incubations, increases in headspace pressure were routinely monitored. Parent compound depletion and formation of methane were confirmed by gas chromatography (GC). The net amount of sulfate and nitrate depletion over the controls was monitored by high pressure liquid chromatrography (HPLC).

Diisopropyl ether is not anaerobically degraded under nitrate- or sulfate-

reducing conditions, and it is not anaerobically degraded under

methanogenic conditions.

Reliability : (2) valid with restrictions

The publication reported a well-documented study that meets basic

scientific principles.

24.02.2006 (32)

Type : aerobic

inoculum : other: Gordonia terrae strain IFP 2001 (CNCM Registration No. CTP 1-

1889); isolated from activated sludge taken at an urban waste water

treatment plant

Contact time : 24 hour(s)

Degradation : (±) % after

Result

Conclusion

Deg. product

Method : other: (sealed flasks, shaken)

Year : 2000 GLP : no

Test substance : other TS: diisopropyl ether (CAS No. 108-20-3)

Result : Diisopropyl ether was degraded by 78% over the 24-hour incubation

period.

Comparison of DIPE biodegradation to ETBE: Test Substance Degradation (%)

ETBE 100

The authors indicated concentrations of the test substance in the flasks were quantified by analytical means. The method was not described in the report, but was referenced in an earlier publication by the same workers.

and t-amyl methyl ether (TAME), but was tested on other ethers including

Test condition: The capacity of ethyl t-butyl ether (ETBE)-induced resting cells of the inoculum to degrade diisopropyl ether was tested in sealed flasks. The article focused on biodegradation of ETBE, methyl t-butyl ether (MTBE)

diisopropyl ether (DIPE).

G. terrae IFP 2001 was cultivated on ETBE-supplemented MM medium. After 24 hours incubation, bacteria were harvested by centrifugation (20,000 g for 20 minutes), washed twice in 100 mM Tris-HCl buffer at pH 7.0 and re-suspended in Tris-HCl. The test substance was added to 20-mL cell suspensions in 125-mL sealed flasks. Flasks were incubated for 24 hours at 30 °C with orbital shaking. Initial cell concentration was 0.5 g/L. The test substance was tested at 100 mg/L. Filtered samples were

analyzed at 0-hour and 24-hours.

Conclusion: Diisopropyl ether was degraded by 78% within 24 hours.

Reliability : (2) valid with restrictions

Information on the analytical method was not provided in the report.

07.02.2006 (21)

Id 108-20-3

Date

3.6 BOD5, COD OR BOD5/COD RATIO

3.7 BIOACCUMULATION

Species: other: see remark

Exposure period : at 25 °C

Concentration

BCF : = 2.95

Elimination

Method : other: calculation

Year

GLP : no

Test substance : other TS: Diisopropyl Ether (CAS # 108-20-3)

Remark : A log bioconcentration factor (BCF) of 0.47 is calculated (BCF = 2.95). With

respect to a log Kow = 1.52, which was used to calculate the BCF, diisopropyl ether in the aquatic environment is expected to have a low

bioaccumulation potential.

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated and not measured.

Flag : Critical study for SIDS endpoint

12.12.2005 (14)

Species: other: see remark

Exposure period : at 25 °C

Concentration

BCF : = 14.06

Elimination :

Method : other: calculation

Year :

GLP : no

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Remark : A log bioconcentration factor (BCF) of 1.15 is calculated (BCF = 14.06).

With respect to a log Kow = 2.4, which was used to calculate the BCF, disiopropyl ether in the aquatic environment is expected to have a low

bioaccumulation potential.

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated and not measured.

12.12.2005 (14)

3.8 ADDITIONAL REMARKS

Memo : Biodegradation of diisopropyl ether

Remark: The article reports on a U.S EPA and American Petroleum Institute

workshop (February 1-3, 2000) on biodegradation of methyl tert-butyl ether (MTBE)-contaminated soils and groundwater. MTBE is one of a group of structurally similar compounds commonly called alkyl ether oxygenates (AEO) that are added to reformulated gasoline to reduce carbon monoxide and ozone emissions. Diisopropyl ether (DIPE) is one type of AEO that is used in gasoline along others in this class of chemicals. The workshop focused on the status of the current research and understanding on biodegradation of MTBE and reported relevant information on the biodegradation of DIPE and other AEOs used in reformulated gasoline.

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Pure microbial cultures have been identified and isolated that have demonstrated the capability of utilizing MTBE as a sole carbon and energy source under aerobic conditions (Scow et al., 2000). This isolate, bacterial strain PM1, was studied by Church and Tratnyek (2000) to determine the aerobic degradation pathway of MTBE. In their study, the authors confirmed the mineralization of MTBE and determined the degradation rates of DIPE and other AEOs were of the same order of magnitude as the degradation rates of MTBE (Church and Tratnyek, 2000). Their results suggested that similar enzyme systems were responsible for all of the reactions.

While the majority of research on anaerobic biodegradation of these compounds has been unable to show that MTBE is utilized, a few studies have demonstrated that MTBE and other AEOs may be susceptible to attack under anaerobic conditions. Kropp et al. (2000) studied the anaerobic biodegradation potential of MTBE, DIPE, and other oxygenates in sediment slurries under methanogenic conditions. They found definite evidence in the form of methane and carbon dioxide production to conclude that anaerobic degradation was occurring. The workshop authors concluded that anaerobic biodegradation was a phenomena that was not widespread and extremely difficult for these compounds.

07.02.2006 (7) (12) (24) (36)

Memo

Biodegradation of diisopropyl ether under aerobic and anaerobic conditions - summary

Remark

: Diisopropyl ether (DIPE) is one of a group of similar compounds referred to as alkyl ether oxygenates (AEO) that are added to reformulated gasoline. Biodegradation studies with DIPE and other AEO compounds have demonstrated the ability of these substances to be consumed by pure strains and mixed cultures of bacteria when incubated under aerobic conditions. Church and Tratnyek (2000) showed that bacterial strain PM1, which had been acclimated to methyl t-butyl ether (MTBE), could mineralize MTBE and demonstrated similar degradation rates for DIPE and other AEOs. They concluded that similar enzymes were responsible for all the degradation reactions. Additional evidence showing the wide spectrum of activity of the bacterial enzyme systems to degrade AEOs was provided by Hernandez-Perez et al. (2001). Using isolated Gordonia terrae (strain IFP 2001) that had been grown on ethyl t-butyl ether (ETBE), degradation of a variety of other AEOs could be achieved. DIPE was degraded 78% within 24 hours in their study (Hernandez-Perez et al. 2001).

Optimum biodegradation in mixed culture systems occurred when the microbial culture is allowed a period of acclimation to the substrate. For example, Bridié et al. (1979) measured only 7% consumption of the theoretical oxygen demand when DIPE was tested in a 5-day BOD test. In contrast. Cano et al. (1999) showed rapid utilization of DIPE when activated sludge was conditioned to a cocktail of volatile organic compounds for two months. In a continuous flow reactor, DIPE removal averaged 99.4%. Cano et al. (1999) also measured high rates of biodegradation of DIPE when comparing the continuous treatment method (EPA Method 304B) (EPA, 1994) to two batch treatment methods (BOX and SBT methods; Rajagopalan et al., 1998). Based on the measured rate constants, the authors considered DIPE to be readily biodegradable. Pruden et al. (2001) also observed high removal rates (e.g., 99.9%) of DIPE through a continuous flow reactor system. The performance of the reactors was enhanced when biomass was retained in the reactor, suggesting that a long biomass residence time may be needed for complete mineralization.

Biodegradation of DIPE is not always observed in biodegradation assays.

Id 108-20-3

Date

Zenker et al. (1999) failed to show biodegradation of DIPE over a 1-year period using indigenous microflora in sediment and water from an aquifer that had been previously exposed to MTBE. Given the apparent need of microbial communities for a period of acclimation to DIPE, it is unlikely that DIPE would be considered readily biodegradable in standard guideline studies. However, the evidence shows that DIPE can be inherently degraded by pure strains of bacteria and mixed enrichments of activated sludge microorganisms.

While available research shows that DIPE is capable of being biodegraded under aerobic conditions, anaerobic biodegradation is extremely difficult and this substance is considered recalcitrant under those conditions. Suflita et al. (1993) showed no biodegradation of DIPE after 252 days of anaerobic incubation. Substrate was added as 50 ppm C to sediment and groundwater collected from a methanogenic portion of a shallow anoxic aguifer. Similarly, DIPE was evaluated for anaerobic biodegradability under methanogenic conditions as well as sulfate and nitrate-reducing conditions (Mormile et al., 1994). Inocula from three sources (e.g., sediment/groundwater from an aquifer impacted by landfill leachate, sediment/surface water from a river impacted by oil storage, and sediment/surface water from a creek impacted by industrial waste and domestic sewage) were used in separate incubations to assess anaerobic biodegradation in sealed serum bottles. No DIPE biodegradation was measured over incubation periods of 85 days (nitrate-reducing conditions), 180 days (methanogenic conditions), and 244 days (sulfate-reducing conditions). Lack of methane production reported by Suflita and Mormile (1993) does not preclude partial anaerobic biodegradation of DIPE in their studies because only methane was monitored. Kropp et al. (2000) studied the anaerobic biodegradation potential of a number of AEOs including DIPE in sediment slurries under methanogenic conditions. They found evidence in the form of methane and carbon dioxide production to conclude that anaerobic biodegradation was occurring, although the authors stated that anaerobic biodegradation was not a widespread phenomena and extremely difficult for these compounds.

27.02.2006

Id 108-20-3 4. Ecotoxicity

Date

ACUTE/PROLONGED TOXICITY TO FISH

flow through Type

Species Pimephales promelas (Fish, fresh water)

Exposure period 96 hour(s) Unit mg/l LC50 = 91.7

Limit test

Analytical monitoring yes

other: Flow-through Fish Acute Toxicity Test Method

Year 1983 **GLP** no data

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method The water solubility of the test chemical was obtained from literature or

> determined experimentally. A flow through system using proportional diluters and modified continuous mini-diluter system was used for

maintaining the required test concentrations

Twenty to twenty-five 30 day-old fish, each weighing approximately 0.12 g, were randomly divided amongst the test tanks (control and five different

concentrations) with flow-through dilutor systems.

Lake Superior water maintained at 25°C ± 1°C was used in the test. Routine measures of hardness (EDTA) and total alkalinity of test water yielded mean values of 45.5 and 42.2 mg/L as CaCO3, respectively. The arithmetic mean of the pH was 7.5 and dissolved oxygen was always

greater than 60% of saturation.

Fish were supplied from the United States Environmental Protection Agency, Environmental Research Laboratory-Duluth culture. They were not fed during the test. Deaths were recorded after 1, 3, 6,12, 24, 48, 72, and

Remark Statistics: Trimmed Spearman-Karber Method

Test method described in reference.

Result 96-hour LL50 = 91.7 mg/L based upon measured values

Analytical method used was GC analysis with Flame Ionization Detection

(GC-FID), performed on a Hewlett-Packard model 5730A gas

chromatograph. Concentrations of the test chemical were measured daily

at each exposure level.

Conclusion 96-hour LC50 = 91.7 mg/L based upon measured values.

Reliability (2) valid with restrictions

This robust summary has a reliability rating of 2 because complete information on the analytical results were not available and the study was

not conducted under GLP.

Critical study for SIDS endpoint Flag

01.11.2005 (43)

Type

Species other: Fish Exposure period 96 hour(s) Unit ma/l LC50 = 214.1

Method other: ECOSAR version 0.99h, US EPA

Year

GLP

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3) 4. Ecotoxicity Id 108-20-3
Date 27.02.2006

Method

: ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs) presented in this program are used to predict the aquatic toxicity of chemicals based on their similarity of structure to chemicals for which the aquatic toxicity has been previously measured. Most SAR calculations in the ECOSAR Class Program are based upon the octanol/water partition coefficient (Kow). SARs have been used by the U.S. Environmental Protection Agency since 1981 to predict the aquatic toxicity of new industrial chemicals in the absence of test data. SARs are developed for chemical classes based on measured test data that have been submitted by industry or they are developed by other sources for chemicals with similar structures, e.g., phenols. Using the measured aquatic toxicity values and estimated Kow values, regression equations can be developed for a class of chemicals. Toxicity values for new chemicals may then be calculated by inserting the estimated Kow into the regression equation and correcting the resultant value for the molecular weight of the compound.

To date, over 150 SARs have been developed for more than 50 chemical classes. These chemical classes range from the very large, e.g., neutral organics, to the very small, e.g., aromatic diazoniums. Some chemical classes have only one SAR, such as acid chlorides, for which only a fish 96-hour LC50 has been developed. The class with the greatest number of SARs is the neutral organics, which has SARs ranging from acute and chronic SARs for fish to a 14-day LC50 for earthworms in artificial soil. The ECOSAR Class Program is a computerized version of the ECOSAR analysis procedures as currently practiced by the Office of Pollution Prevention and Toxics (OPPT). It has been developed within the regulatory constraints of the Toxic Substances Control Act (TSCA). It is a pragmatic approach to SAR as opposed to a theoretical approach.

Result : LC50, 96 h, for fish = 214.1 mg/L

Test condition : Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al, 1987),

log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C (SRC

PhysProp database) were entered into the program.

Class: Neutral organics

Test substance: Diisopropyl Ether (CAS No. 108-20-3)

Conclusion : The predicted 96 h LC50 value for fish

The predicted 96 h LC50 value for fish (214.1 mg/L) is in good agreement with the experimental 96 h LC50 value for fathead minnow (Pimephales promelas) (91.7 mg/L) (Veith et al., Can. J. Fish. Aquat. Sci., 40:743-748) and 48 h EC50 value for Daphnia (190.0 mg/L) (Stephenson R.R., Shell

Research Limited, Report No. SBGR.83.215).

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated and not measured.

28.10.2005 (13)

Type : flow through

Species: Pimephales promelas (Fish, fresh water)

Exposure period : 96 hour(s)
Unit : mg/l
LC50 : = 786
EC50 : = 476

Limit test

Analytical monitoring : yes

Method : other: Flow-through Fish Acute Toxicity Test

Year : 1983 GLP : no data

Test substance: other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method : Test solutions were prepared using a proportional diluter system without

replication. This system provided control and five test substance concentrations to glass test vessels. Each vessel held 2 L of test solution and the diluter flow rate was sufficient to provide 18 volume additions per day. An aqueous stock solution of 1050 mg/L was used by the diluter to

4. Ecotoxicity Id 108-20-3
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prepare the exposure series. Dilution water was filtered Lake Superior water. Typical ranges of water quality factors measured in this water were pH (7.4 - 8.2), total hardness (44 - 53 mg/L as CaCO3), and specific conductance (78 - 86 mmhos/cm).

Test fish originated from in-house cultures of P. promelas at the U.S. EPA Environmental Research Laboratory - Duluth. Fish were not fed 24 h prior to testing or during the test. At test initiation, fish were randomly placed in test vessels until each vessel contained 10 individuals. Individuals used in testing were 34 d old and measured 19.0 mm mean length (SD = 2.534) and 0.104 g mean weight (SD = 0.0433). Biomass loading was 1.04 g/L. Death was the major test endpoint. Numbers of dead fish were counted daily and any dead fish were removed from the vessels. Abnormal behavioral changes were recorded at each observation time. LC50 (lethality) and EC50 (total effect) values were determined.

Temperature, dissolved oxygen, and pH were measured daily in all test chambers. Mean values (and Standard Deviation) were 24.9 °C (SD = 0.52), 7.3 mg/L (SD = 0.13), and 7.75 (SD = 0.16), respectively. Total hardness and alkalinity were measured once in the control, low, medium, and high test levels. Mean values were 43.7 mg/L total hardness as CaCO3 (SD = 0.96) and 49.6 mg/L alkalinity as CaCO3 (SD = 0.25). Lighting was provided by fluorescent bulbs that produced 28 to 48 lumens/sq ft at the water surface. The photoperiod was 16 h light and 8 h dark.

Test substance concentrations were verified in most cases daily during the test using gas-liquid chromatography. Concentrations were averaged and a mean percent recovery was calculated. The nominal with measured concentrations in parentheses were, control (not detected), 157 mg/L (131 mg/L), 242 mg/L (210 mg/L), 373 mg/L (382 mg/L), 574 mg/L (594 mg/L), and 883 mg/L (1044 mg/L). The overall percent recovery was 98.7%.

Remark: Statistics: LC/EC50 values determined by Trimmed Spearman-Karber

Method

Result : 96-hour LC50 = 786 mg/L based on mean measured values.

96-hour EC50 = 476 mg/L based on mean measured values.

The EC50 value was based on mortality and the following abnormal effects: loss of schooling behavior, swimming near the surface, hypoactive,

under-reactive to external stimuli, loss of equilibrium.

Conclusion : 96-hour LC50 = 786 mg/L based on mean measured values.

96-hour EC50 = 476 mg/L based on mean measured values.

Reliability : (1) valid without restriction

12.12.2005 (16)

Type : flow through

Species: Pimephales promelas (Fish, fresh water)

Exposure period : 96 hour(s) **Unit** : mg/l **LC50** : = 900

Limit test

Analytical monitoring : yes

Method : other: Flow-through Fish Acute Toxicity Test (ASTM, 1980)

Year : 1985 GLP : no data

Test substance: other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method: Test solutions were prepared using a continuous-flow diluter delivery

system, which delivered four test substance concentrations and control solutions to duplicate test vessels. Dilution water was filtered Lake Superior water. Average values for water quality factors for the dilution water were: hardness (44.6 mg/L as CaCO3), total alkalinity (44.0 mg/L as

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> CaCO3), and pH (7.6). Test chambers were glass vessels and contained 2 L of test solution. Solution flow rates through the test chambers was sufficient to provided at least a 95% replacement in approximately 4 h. Test substance concentrations were verified daily during the test using either gas chromatography or high pressure liquid chromatography methods.

> The mean temperature for the test was 25 ± 0.5 °C, and dissolved oxygen remained at or above 80% saturation. Lighting was provided by wide spectrum fluorescent bulbs at an intensity of 22 to 38 lumens/sq ft over the test chambers. The photoperiod was 16 h light and 8 h dark with a 30-min dusk/dawn transition period.

> Test fish originated from cultures maintained by the U.S. EPA Environmental Research Laboratory - Duluth, MN. and were 28 to 34 days old (weighing approximately 0.12 g) at the time of testing. A total of 20 fish per treatment (10/replicate) was used in the test. Fish were added to the test chambers 2-3 h before introduction of the test solutions. Fish were not fed 24 h before or during the test. Mortalities were recorded daily.

Statistics: Trimmed Spearman-Karber Method or log-probit method. Remark

Result 96-h LC50 = 900 mg/L based on measured concentrations

95% CL = 881 - 920 mg/L

96-h LC50 = 900 mg/L based on measured concentrations Conclusion

(1) valid without restriction Reliability

12.12.2005 (3)

Type static

Species Carassius auratus (Fish, fresh water)

Exposure period 24 hour(s) Unit mg/l LC50 = 380

Limit test

Analytical monitoring yes

Method other: static acute fish toxicity test (APHA, 1971)

Year

GLP

Test substance other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method The test consisted of exposing groups of six fish to a series of

> concentrations of the test substance for 24 h. Fish were exposed in an all glass tanks holding 25 liters of test solution. Dilution water was local tap

water having the following characteristics (all values in mg/L):

 $CI^{-} = 65$; $NO2^{-} = 0$; $NO3^{-} = 4$; $SO4^{-}2 = 35$; $PO4^{-}3 = 0.15$; $HCO3^{-} = 25$; SiO2 = 25; NH4+ = 0; Fe = 0.05; Mn = 0; Ca+2 = 100; Mg+2 = 8; alkali as

Na+ = 30; pH = 7.8.

The test was run at a temperature of 20±1°C, and the solutions were not aerated during the test period.

Test fish had a mean length of 6.2 ± 0.7 cm, a mean weight of 3.3 ± 1.0 g and were in good health at the time of testing.

Exposure concentrations were confirmed either by total organic carbon analysis or by extraction and subsequent analysis by gas chromatography.

Measured concentrations were not reported in this study.

Remark Determination of LC50 by graphical interpolation of log concentrations

versus percent mortality (APHA, 1971).

Result 24-hour LC50 = 380 mg/L

> The analytical method was either total organic carbon analysis or gas chromatography. It was not reported what method was employed for this test substance nor if the result was based on measured concentrations.

Date

Conclusion : 24-hour LC50 = 380 mg/L.

Reliability : (3) invalid

The test was run for only 24 hours to ensure that the dissolved oxygen content did not fall below 4 mg/L. The report lacked sufficient detail for assessment. It was not stated whether results were based on nominal or

measured values.

12.12.2005 (1)

Type : static

Species: Lepomis macrochirus (Fish, fresh water)

 Exposure period
 : 96 hour(s)

 Unit
 : mg/l

 LC50
 : 7000

Limit test

Analytical monitoring : no

Method : other: static acute fish toxicity test

Year :

GLP : no

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method: The test consisted of exposing groups of fish to a four-dilution series of the

test substance for a period of 96 h. Test vessels were all glass 5-gallon aquaria. The volume of test solution was adjusted to assure that a biomass loading was no more than 1 g fish /liter solution. Dilution water was well water having a typical pH of 7.6 to 7.9 and a hardness of 55 mg/L

(as CaCO3).

Fish were obtained from a commercial source and assessed for health during a 14-d acclimation period prior to testing. During that time they were maintained on a commercial fish food diet supplemented with minced frozen shrimp. Fish were not fed 48 hours prior to testing. Fish were randomly selected for testing and were approximately 33 to 75 mm in length.

The test was run at 23°C. Test solutions were not aerated for the initial 24 h, but aeration was applied thereafter if the dissolved oxygen concentration was being depleted. Dissolved oxygen readings were taken daily, and pH was measured at the end of the test. However, these data were not provided in the report.

Mortality was assessed daily and any dead fish were removed at each

observation time.

Remark: The LC50 was determined by plotting survival percentages on semi-

logarithmic paper and drawing a straight line fit through or near significant

points above and below 50% survival.

Result : 96-hour LC50 = 7,000 mg/L

The mortality pattern reported for the test substance suggests that a more likely estimate of the LC50 value would lie between 7,900 and 10,000 mg/L, rather than 7,000 mg/L. This was due to a non-monotonic dose response pattern of mortality. The report authors indicated that the LC50 value was higher than the published solubility for the test substance.

Conclusion : 96-hour LC50 = 7,000 mg/L based on nominal concentrations

Reliability : (3) invalid

Documentation was insufficient for evaluation. Basic water quality data during the test were not provided. The authors stated that aeration of the test solutions was used after 24 hours to ensure maintenance of dissolved oxygen. No analytical verification of exposure concentrations were made.

12.12.2005 (10)

Type : static

Species: Menidia beryllina (Fish, estuary, marine)

Date

 Exposure period
 : 96 hour(s)

 Unit
 : mg/l

 LC50
 : 6600

Limit test

Analytical monitoring : no

Method : other: static acute fish toxicity test

Year :

GLP : no

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method: The test consisted of exposing groups of fish to a four-dilution series of the

test substance for a period of 96 h. Test vessels were all glass 5-gallon aquaria. The volume of test solution was adjusted to assure that a biomass loading was no more than 1 g fish /liter solution. Dilution water was prepared by adding "instant ocean" salts to well water (pH of 7.6 to 7.9; hardness 55 mg/L (as CaCO3)) until a specific gravity of 1.018 was

achieved.

Fish were field collected in nets from Horseshoe Bay at Sandy Hook, New Jersey. They were held for a 14-d acclimation period prior to testing and assessed for health during that time. During the acclimation period they were fed minced frozen shrimp. Fish were not fed 48 hours prior to testing. Fish were randomly selected for testing and were approximately 40 to 100 mm in length.

The test was run at 20°C, and test solutions were continuously aerated during the exposure period. Dissolved oxygen readings were taken daily, and pH was measured at the end of the test. However, these data were not provided in the report.

Mortality was assessed daily and any dead fish were removed at each

observation time.

Remark : LC50 determined by graphical interpolation of the logarithm of the

concentration versus the percentage mortality.

Result : 96-hour LC50 = 6600 mg/L

The mortality pattern reported for the test substance does not correspond with the estimated LC50 value. Given the dose-response pattern, the LC50 value would lie between 3,200 and 5,000 mg/L, rather than 6,600 mg/L. The authors reported that the result was higher than the reported

water solubility of the test substance.

Conclusion: 96-hour LC50 = 6600 mg/L based on nominal concentrations.

Reliability : (3) invalid

Documentation was insufficient for evaluation. Basic water quality data during the test were not provided. The authors stated that aeration of the test solutions was used after 24 hours to ensure maintenance of dissolved oxygen. No analytical verification of exposure concentrations were made.

12.12.2005 (10)

4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

Type :

Species : Daphnia magna (Crustacea)

Exposure period : 48 hour(s)
Unit : mg/l
EC50 : = 190
Analytical monitoring : no

Method : other: U.S. Environmental Protection Agency, Methods for acute toxicity

testing with fish, macro-invertebrates and amphibians (EPA-660/3-75-009)

Year : 1975

Date

GLP : no

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Remark : Statistics:

Probit analysis after log transformation of the concentrations (Finney, 1971) Probit Analysis, Finney, D.J., Cambridge University Press, 3rd edition,

p333 (1971)

Result : The 24 h and 48 h Effect Concentration (EC50) values were calculated to

be 240 mg/L (95% fiducial limits 210 to 280 mg/L) and 190 mg/L (95%

fiducial limits 160 to 220 mg/L), respectively.

The immobilization (%) of Daphnia magna (n=10/replicate) are as follows:

Test Substance Immobilization (%)*

Loading Rate

(mg/L) 24 hr 48 hr

0 (control)	0	0
46	0	0
99	3	7
210	27	57
460	100	100
1000	100	100

*mean of 3 replicates

Test condition : A 48 hour static toxicity test was carried out without renewal of the test

solutions. Quantities of stock solutions of di-isopropyl ether in acetone were added in triplicate sets of 110 mL glass flasks so that when made up with

reconstituted freshwater, an approximately logarithmic series of

concentrations ranging from 46 to 1000 mg/L was produced. Three flasks served as controls and received no test substance. The concentration of acetone in all control and test flasks was 0.1 mL/L. Precautions were taken to (a) minimise evaporative loss of the test substance by use of glass cover slips over the vessel necks and (b) to minimize the risk of organisms becoming trapped at the surface by placing black paper caps over the flasks to create a darkened zone which the organisms would avoid.

The test temperatures were in the range $20 \pm 2^{\circ}$ C, pH was in the range 8.2 to 8.4, the total hardness was 164 mg/L as CaCO3, and dissolved oxygen was in the range 8.2 to 9.2 mg/L.

The daphnids were cultured in-house, derived from a strain obtained (via ICI Brixham Laboratory) from Institut National de Recherche Chimique Applique (I.R.Ch.A.), France. Age was <24 hours old from 15 to 35 day old parents.

All concentrations of test substance are expressed in terms of quantities

initially added to the test vessels.

Test substance : Diisopropyl Ether (CAS No. 108-20-3)

Conclusion : After Daphnia magna were exposed to test solutions of di-isopropyl ether

for 48 hours in a static test, the 24 h and 48 h EC50 values were calculated

to be 240 mg/L and 190 mg/L, respectively.

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because it did not analytically verify exposure concentrations and the results are based on

nominal values.

07.12.2005 (38)

Type

Species : other: Daphnia
Exposure period : 48 hour(s)
Unit : mg/l
EC50 : = 221.9

Method : other: ECOSAR version 0.99h, US EPA

Year

GLP :

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Date

Method

: ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs) presented in this program are used to predict the aquatic toxicity of chemicals based on their similarity of structure to chemicals for which the aquatic toxicity has been previously measured. Most SAR calculations in the ECOSAR Class Program are based upon the octanol/water partition coefficient (Kow). SARs have been used by the U.S. Environmental Protection Agency since 1981 to predict the aquatic toxicity of new industrial chemicals in the absence of test data. SARs are developed for chemical classes based on measured test data that have been submitted by industry or they are developed by other sources for chemicals with similar structures, e.g., phenols. Using the measured aquatic toxicity values and estimated Kow values, regression equations can be developed for a class of chemicals. Toxicity values for new chemicals may then be calculated by inserting the estimated Kow into the regression equation and correcting the resultant value for the molecular weight of the compound.

To date, over 150 SARs have been developed for more than 50 chemical classes. These chemical classes range from the very large, e.g., neutral organics, to the very small, e.g., aromatic diazoniums. Some chemical classes have only one SAR, such as acid chlorides, for which only a fish 96-hour LC50 has been developed. The class with the greatest number of SARs is the neutral organics, which has SARs ranging from acute and chronic SARs for fish to a 14-day LC50 for earthworms in artificial soil. The ECOSAR Class Program is a computerized version of the ECOSAR analysis procedures as currently practiced by the Office of Pollution Prevention and Toxics (OPPT). It has been developed within the regulatory constraints of the Toxic Substances Control Act (TSCA). It is a pragmatic approach to SAR as opposed to a theoretical approach.

Result : EC50, 48 h, for Daphnia = 221.9 mg/L

Test condition : Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al, 1987),

log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C (SRC

PhysProp database) were entered into the program.

Class: Neutral organics

Test substance: Diisopropyl Ether, CAS No. 108-20-3

Conclusion : The predicted 48 h LC50 value for Daphnia (221.9 mg/L) is in close

agreement with the experimental 48 h EC50 value for Daphnia (190.0

mg/L) (Stephenson R.R., Shell Research Limited, Report No.

SBGR.83.215).

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated and not measured.

28.10.2005 (13)

4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE

Species : other algae: Green Alga

Endpoint

Exposure period : 96 hour(s)
Unit : mg/l
EC50 : = 134.9
ChV : = 10.2

Method : other: ECOSAR version 0.99h, US EPA

Year

GLP :

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method: ECOSAR version 0.99h, US EPA. The structure-activity relationships

(SARs) presented in this program are used to predict the aquatic toxicity of chemicals based on their similarity of structure to chemicals for which the

Date

aquatic toxicity has been previously measured. Most SAR calculations in the ECOSAR Class Program are based upon the octanol/water partition coefficient (Kow). SARs have been used by the U.S. Environmental Protection Agency since 1981 to predict the aquatic toxicity of new industrial chemicals in the absence of test data. SARs are developed for chemical classes based on measured test data that have been submitted by industry or they are developed by other sources for chemicals with similar structures, e.g., phenols. Using the measured aquatic toxicity values and estimated Kow values, regression equations can be developed for a class of chemicals. Toxicity values for new chemicals may then be calculated by inserting the estimated Kow into the regression equation and correcting the resultant value for the molecular weight of the compound.

To date, over 150 SARs have been developed for more than 50 chemical classes. These chemical classes range from the very large, e.g., neutral organics, to the very small, e.g., aromatic diazoniums. Some chemical classes have only one SAR, such as acid chlorides, for which only a fish 96-hour LC50 has been developed. The class with the greatest number of SARs is the neutral organics, which has SARs ranging from acute and chronic SARs for fish to a 14-day LC50 for earthworms in artificial soil. The ECOSAR Class Program is a computerized version of the ECOSAR analysis procedures as currently practiced by the Office of Pollution Prevention and Toxics (OPPT). It has been developed within the regulatory constraints of the Toxic Substances Control Act (TSCA). It is a pragmatic approach to SAR as opposed to a theoretical approach.

Result : EC50, 96 h, for green algae = 134.9 mg/L

ChV, 96 h, for green algae = 10.2 mg/L

Test condition : Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al, 1987),

log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C (SRC

PhysProp database) were entered into the program.

Class: Neutral organics

Test substance: Diisopropyl Ether (CAS No. 108-20-3)

Conclusion : The predicted 96 h EC50 value for algae (134.9 mg/L) is in the same range

as the predicted 48 h LC50 value for Daphnia (221.9 mg/L) and the predicted 96 h LC50 value for fish (214.1 mg/L). There is also good comparison between the predicted and experimental EC50 values for Daphnia (221.9 mg/l v 190.0 mg/L, respectively) and for fish (214.1 mg/l v

91.7 mg/L, respectively).

Reliability : (2) valid with restrictions

This robust summary has a reliability rating of 2 because the data are

calculated and not measured.

28.10.2005 (13)

Species : Selenastrum capricornutum (Algae)

 Endpoint
 : biomass

 Exposure period
 : 96 hour(s)

 Unit
 : mg/l

 EC50
 : >= 1000

Limit test :

Analytical monitoring : no

Method : other: algae growth inhibition

Year : 1983 GLP : no data

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method : A 4 d algal growth study was carried out using 10 concentrations of the test

substance and a control. The test design included six control replicates and single vessels dosed with different concentrations of the test

substance. 250-mL glass Erlenmeyer flasks served as the test vessels and held 50 mL of culture medium. Culture medium was prepare following the recipe given by Miller and Green (1978) with the following exceptions; 1)

Date

boric acid concentration = 105 mg/L, and 2) sodium bicarbonate concentration = 50 mg/L.

To 10 flasks, quantities of a test substance stock solution made up in acetone were added to give a logarithmic series of concentrations ranging from 1 to 1000 mg/L (1.0, 2.2, 4.6, 10, 22, 46, 100, 220, 460, and 1000 mg/L). The concentration of acetone in all flasks including controls was adjusted to 0.1 mL/L. Each flask was inoculated with S. capricornutum to give an initial cell density of 5 x 102 cells/mL. The algal inoculum was prepared from an actively growing liquid culture of S. capricornutum in exponential growth phase.

Flasks were incubated in a temperature controlled orbital incubator under constant illumination (approximately 3000 lux) at 24±2°C for 4 days. Cell counts were made on days 2 and 4 using an electronic particle counter (Coulter counter). The temperature in the incubator was measured at 4-h intervals. The pH of the control and highest test concentration was measured on days 0, 2, and 4. Temperature remained within the 24±2°C specified range, and the pH ranged from 8.3 to 8.5 in the measured vessels.

All determination of EC50 values were based on nominal test concentrations and cell counts.

Result: 96-hour EC50 = >1000 mg/L based on nominal concentrations.

The 96-hour cell counts in the treated flasks as a percent of the mean control cell counts were:

1.0 mg/L = 84% 46 mg/L = 127%

2.2 mg/L = 108% 100 mg/L = 130%

4.6 mg/L = 91%220 mg/L = 113%

10 mg/L = 122% 460 mg/L = 127% 22 mg/L = 129% 1000 mg/L = 91%

Conclusion : 96-hour EC50 = >1000 mg/L based on nominal concentrations. **Reliability** : (3) invalid

Test concentrations were not measured and there is no indication in the report whether the test vessels were sealed. The reported LC50 value may reflect a loss of test substance by volatilization if the flasks were not tightly

sealed.

12.12.2005 (39)

4.4 TOXICITY TO MICROORGANISMS E.G. BACTERIA

4.5.1 CHRONIC TOXICITY TO FISH

4.5.2 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES

4.6.1 TOXICITY TO SEDIMENT DWELLING ORGANISMS

4.6.2 TOXICITY TO TERRESTRIAL PLANTS

4.6.3 TOXICITY TO SOIL DWELLING ORGANISMS

4. Ecotoxicity		ld 108-20-3
	Da	te 27.02.2006
4.6.4 TOX. TO OTHER NON MAMM. TE	RR. SPECIES	
4.7 BIOLOGICAL EFFECTS MONITOR	RING	
4.8 BIOTRANSFORMATION AND KIN	ETICS	
4.9 ADDITIONAL REMARKS		
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5. Toxicity Id 108-20-3

Date

5.0 TOXICOKINETICS, METABOLISM AND DISTRIBUTION

5.1.1 ACUTE ORAL TOXICITY

Type : LD50

Value

Species : rat

Strain : Sprague-Dawley
Sex : male/female

Number of animals

Result

Vehicle : other: None; administered undiluted

Doses :

Method : other: Similar to OECD 401

Year

GLP : no

Test substance : other TS: Diisopropyl ether (CAS No. 108-20-3)

Method : Administered orally to nonfasted rats. LD50 calculated by the method of

Litchfield and Wilcoxon [1949]. Similar to OECD 401.

Remark : Test type: Acute oral toxicity

Year: Prior to 1971

No. of animals/dose: 6 male for young adult and older adult

6 - 12 male and female for 14-day old rats Route of administration: Oral gavage

Dose level: Variable Dose volume: Variable

Control group included: No, but none needed 14-day old: LD50 6.4 ml/kg [approx 4.5 g/kg]

young adults: LD50 16.5 ml/kg [approx 11.6 g/kg] Older adults: LD50 16.0 ml/kg [approx 11.2 g/kg]

G/kg dose based on a density of 0.72 g/ml

Test condition : Rats were observed for up to 7 days after dosing.

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis [propane],

2,2 '-

Source/purity of test material is not specified, but stated to be analytical

grade meeting ACS specifications.

Conclusion : DIPE, when administered to adult male Sprague-Dawley rats, had an acute

oral LD50 of >10 g/kg. 14-day immature rats were considerable more

sensitive [LD50 4.5 g/kg].

Reliability : (2) valid with restrictions

Not GLP but conducted at a reputable laboratory [Abbot Laboratories,

Chicago].

01.11.2005 (23)

Type : Value :

Species : rabbit

Strain : New Zealand white

Sex : no data Number of animals : 6

Vehicle : other: none reported

Doses : 8.2, 7.2, 6.0, 5.2, 3.3, 1.62 g/kg **Method** : other: Similar to OECD 401

Year :

GLP : no

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Test substance : other TS: Diisopropyl ether (CAS No. 108-20-3)

Remark : Test type: Acute oral toxicity

Year: Prior to 1939

Route of administration: Oral

Dose levels: 8.2, 7.2, 6.0, 5.2, 3.3, 1.62 g/kg

Dose volume: Variable Control: No - none needed

Result : Minimal lethal dose between 7 - 9 ml/kg

The symptoms noted were lack of coordination and unsteadiness at onset followed by a slight narcosis. In the animals that died the narcosis progressed towards a deep narcosis with loss of corneal reflex and evidences of depressant action on the medulla appeared, respiration became progressively slower, irregular and variable in amplitude and drop in body temperature till respiration failed. In the surviving animals, no effect on HB, erythrocyte count, total and differential leukocyte count was

observed. No delayed toxicity was observed during the recovery period of 4

months after treatment.

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis [propane],

2,2 '-

Test material is stated to be commercial grade with 3% of isopropyl

alcohol, but with no peroxide nor added inhibitor.

Conclusion: The test article, when administered orally as received to New Zealand

white rabbits had a minimal lethal dose of 7 - 9 ml/kg [approx 5 - 6.5 g/kg].

Reliability : (2) valid with restrictions

Not conducted by GLP but at a reputable laboratory [Kettering Laboratory,

University of Cincinnati].

01.11.2005 (26)

5.1.2 ACUTE INHALATION TOXICITY

Type : Value :

Species : quinea pig

Strain : other: not specified

Sex : no data

Number of animals

Vehicle : other: none

Doses : 0.3%; 1%; 3%; 6% in air

Exposure time

Method : other: not specified

Year :

GLP : no

Test substance : other TS: Diisopropyl ether (CAS No. 108-20-3)

Remark : Test type: Acute inhalation toxicity

Year: Prior to 1939

No. animlas/sex/group: One to two animals per dose

Route of administration: Inhalation Dose level: 0.3%; 1%; 3%; 6% in air

Dose volume: N/A Control: No

Result : 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action

1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia

6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr

Test condition : 1 or 2 hrs or until death [6%]

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis [propane],

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2.2 '-

Test material is stated to be commercial grade with 3% of isopropyl

alcohol, but with no peroxide or added inhibitor.

Conclusion: The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in

three species.

Reliability : (2) valid with restrictions

Not conducted by GLP. Few animals per group, but at a reputable

laboratory [Kettering Laboratory, University of Cincinnati].

01.11.2005 (26)

Type : Value :

Species : rabbit

Strain : New Zealand white

Sex : no data

Number of animals

Vehicle : other: none

Doses : 0.3%; 1%; 3%; 6% in air

Exposure time

Method : other: not specified

Year :

GLP : no

Test substance : other TS: Diisopropyl ether (CAS No. 108-20-3)

Remark: Test type: Acute inhalation toxicity

Year: Prior to 1939

No. animlas/sex/group: One to two animals per dose

Route of administration: Inhalation Dose level: 0.3%; 1%; 3%; 6% in air

Dose volume: N/A Control: No

Result : 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action

1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia

6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr

Test condition : 1 or 2 hrs or until death [6%]

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis [propane],

2,2 '-

Test material is stated to be commercial grade with 3% of isopropyl

alcohol, but with no peroxide or added inhibitor.

Conclusion: The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in

three species.

Reliability : (2) valid with restrictions

Not conducted by GLP. Few animals per group, but at a reputable

laboratory [Kettering Laboratory, University of Cincinnati].

01.11.2005 (26)

Type : Value :

Species : monkey

Strain : other: Macacus rhesus

Sex : female

Number of animals

Vehicle : other: none

Doses : 0.3%; 1%; 3%; 6% in air

Exposure time

Method : other: not specified

Year :

GLP : no

Test substance : other TS: Diisopropyl ether (CAS No. 108-20-3)

Remark : Test type: Acute inhalation toxicity

5. Toxicity Id 108-20-3

Date

Year: Prior to 1939

No. animlas/sex/group: One to two animals per dose

Route of administration: Inhalation Dose level: 0.3%; 1%; 3%; 6% in air

Dose volume: N/A Control: No

Result : 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action

1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia

6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr

Test condition : 1 or 2 hrs or until death [6%]

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis [propane],

2,2 '-

Test material is stated to be commercial grade with 3% of isopropyl

alcohol, but with no peroxide or added inhibitor.

Conclusion : The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in

three species.

Reliability : (2) valid with restrictions

Not conducted by GLP. Few animals per group, but at a reputable

laboratory [Kettering Laboratory, University of Cincinnati].

01.11.2005 (26)

5.1.3 ACUTE DERMAL TOXICITY

Type : LD50

Value :

Species : rabbit

Strain : New Zealand white

Sex : no data

Number of animals

Vehicle : other: none Doses : variable

Method : other: Similar to OECD 402

Year :

GLP : no

Test substance : other TS: Diisopropyl ether (CAS No. 108-20-3)

Remark: Test type: Acute dermal toxicity

Year: Prior to 1939

No. of animals/sex/group: Unspecified Route of administration: Dermal

Dose level: variable

Control: No

Result: No deaths or systemic effects were reported. In rabbits dermal unoccluded

LD50 > 2.0 g/kg. The actual dose applied was much higher, but continued

to evaporate from the skin during application.

Test condition: The material was continuously dripped onto the shaved skin to keep it wet

for one hour, while continuously evaporating. 150 ml of material was used.

Test substance : Diisopropyl ether (CAS No. 108-20-3)

Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis

[propane], 2,2 '-

Test material is stated to be commercial grade with 3% of isopropyl

alcohol, but with no peroxide or added inhibitor.

Conclusion: The test article, when administered dermally to New Zealand white rabbits

had an acute dermal LD50 of greater than 2.0 g/kg.

Reliability : (2) valid with restrictions

Not GLP but conducted at a reputable laboratory [Kettering Laboratory,

University of Cincinnati].

01.11.2005 (26)

5. Toxicity Id 108-20-3

Date

5.1.4 ACUTE TOXICITY, OTHER ROUTES

5.2.1 SKIN IRRITATION

5.2.2 EYE IRRITATION

5.3 SENSITIZATION

Type : other: In vitro chemical reactivity assay, surrogate for respiratory

sensitization

Species: other: No animals; in vitro chemical assay

Number of animals : 0

Vehicle: other: NoneResult: not sensitizingClassification: not sensitizing

Method : other: No guideline available

Year : 1990 **GLP** : no

Test substance: other TS: Diisopropyl ether (CAS No.108-20-3)

Remark: Route of administration: N/A

Sex: N/A
Dose level: N/A
Dose volume: N/A

Control group included: Positive and negative controls included

Result : Diisopropanol was negative in this in vitro assay for potential respiratory

sensitization. The assay gave positive responses with several known

respiratory sensitizers.

Test condition : A method for monitoring chemical reactivity in aqueous solutions, at neutral

pH and 37 degrees C, was developed. The chemical was allowed to react with a lysine-containing peptide, and the reaction was monitored with high-performance liquid chromatography. Simple acids, bases, and solvents did

not react with the peptide, whereas isocyanates, anhydrides, and chloramine-T, substances well known for their sensitizing and asthma inducing properties, did. Thus a positive test strongly suggested that the chemical had the potential to act as a hapten and cause sensitization when

inhaled.

Test substance: Diisopropyl ether (CAS No.108-20-3)

Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis

[propane], 2,2 '-

Source/purity not specified.

Conclusion: Di-isopropanol was negative in this in vitro assay.

Reliability : (2) valid with restrictions

Not conducted by GLP; research method not accepted by regulatory

agencies; in vitro surrogate for respiratory sensitisation.

01.11.2005 (44)

5.4 REPEATED DOSE TOXICITY

Type : Sub-chronic

Species : rat

Sex : male/female
Strain : Sprague-Dawley
Route of admin. : inhalation

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Exposure period : 6 hours/day

Frequency of treatm. : 5 days/week for ~13 weeks

Post exposure period

Doses : 0, 480, 3300, or 7100 ppm

Control group : other: yes (untreated & sham-exposed)

NOAEL : = 480 ppm

Method : EPA OTS 798.2450

Year : 1996 GLP : no data

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method : Statistical method:

Statistical analyses of numerical data included ANOVA and Tukey's studentized range test for data on serum chemistry. Duncan's multiple range test was used for hematology and body weights to assess statistically significant differences between control and exposed groups.

Remark : Male and female rats were acclimated for 2 weeks before initiation of

exposures that began at ~8 weeks of age. Exposed animals were individually housed in 1-m3 inhalation chambers. Untreated control animals were housed in a separate room in identical caging. Room environment was set to 20-22°C and 40-60% relative humidity. Lights were on a 12/12-hr light/dark cycle. Food and water were provided ad libitum except during

exposures.

Vapors were generated by metering DIPE on to a warmed fiberglass wick and carried to the three 1m3 exposure chambers by a stream of measured and filtered room air. Temperature and humidity in the chambers were measured every 30 minutes and air flow through each chamber was metered to provide at least 12 air changes/hour. Chamber air samples were collected with a gas-tight syringe and analyzed with a gas chromatograph with a flame ionization detector and a fused silica column. Samples were periodically obtained for analysis by gas chromatography/mass spectroscopy.

The primary endpoints during the course of exposures were individual body weights recorded weekly and clinical signs recorded daily except on weekends.

Following the last exposure, rats were fasted overnight and weighed; blood samples were obtained via the orbital sinus using light anesthesia. Samples were used to determine values for WBC, RBC, Hgb, Hct, MCV, MCH, MCHC, platelets, and differential counts. Glucose, urea nitrogen, total protein, albumin, globulin, A/G ratio, sorbitol dehydrogenase, aspartate aminotransferase, alanine aminotransferase, alkaline phosphatase, total bilirubin, creatinine, cholesterol, triglycerides, uric acid, Cl, Ca, Na, K, and P. Following the collection of blood samples, all animals were euthanized with sodium pentobarbital (i.p.) and exanguinated. Approximately 40 tissues were collected for histopathology and organs were weighed. Testis and associated tissues were preserved whole in 10% buffered formalin except for the left cauda epididymis of 10 rats in both control groups and the highest test group; epididymides were evaluated for morphology and number of sperm. The left testis in these groups was weighed and used for determination of number of testicular spermatids.

Type: 90-Day Subchronic

Species/strain: Rat/Sprague-Dawley; Tac:N(SD)fBR

No./sex/dose: 14/sex/group

Vehicle: None

Method: USEPA 1984, 40CFR Part 798:2450

Result: DIPE did not adversely affect clinical signs body weight, serum chemistry, hematology, or the number of sperm or spermatids. Exposure to males at

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7100 ppm resulted in hypertrophy of liver cells associated with increased liver weight and in increased kidney weight with an increased incidence of hyaline droplets in proximal tubules of the kidney. Females had increased weight of both liver and kidney, although kidney increased only in relation to sham-exposed controls and no morphologic changes were observed in either organ. At 3300 ppm, weights of liver and kidney were increased in males; the liver weights were increased in females only compared to sham-exposed controls and not untreated controls. No morphologic abnormalities were observed. No changes were observed with 480 ppm.

Test substance: Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.

Conclusion : NOAEL = 480 ppm
Reliability : (2) valid with restrictions

GLP unknown. Study well documented, meets generally accepted scientific

principles, acceptable for assessment.

01.11.2005 (8)

Type : Sub-chronic

Species : rat

Sex: male/femaleStrain: Sprague-DawleyRoute of admin.: inhalationExposure period: 6 hours/day

Frequency of treatm. : 5 days/week for ~13 weeks

Post exposure period :

Doses : 0, 450, 3250, or 7060 ppm **Control group** : other: yes (sham-exposed)

Method : other: USEPA 1984, 40CFR Part 798:6050, 6400, and 6200

Year : 1997 GLP : no data

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method : Statistical method:

All statistical analyses were performed with SAS software. Body weights, rectal temperatures fore- and hindlimb grip strengths, the number of rears, and motor activity were analyzed by a one-way analysis of variance followed by Duncan's multiple range test. The remaining data from the FOB were analyzed by Fisher's exact test using an extended contingency table containing all four groups of at given sex at a specified time. If a significant difference occurred for a given parameter, Fisher's exact test was used to directly compare each group individually against the control. Brain weights,

lengths and widths, were analyzed by Student's t-test.

Remark: Male and female rats were acclimated for 2 weeks before initiation of

exposures that began at ~8 weeks of age. Sham-exposed and exposed animals were individually housed in 1-m3 inhalation chambers except during behavioral testing, when they were placed in another room overnight and evaluated the following day. Room environment was set to 20-22°C and 40-60% relative humidity. Lights were on a 12/12-hr light/dark cycle. Food and water were provided ad libitum except during exposures. Exposure vapors were generated by metering DIPE on to a warmed fiberglass wick and carried to the three 1-m3 exposure chambers by a stream of measured and filtered room air. Temperature and humidity in the chambers were measured every 30 minutes and air flow through each chamber was metered to provide at least 12 air changes/hour. Chamber air samples were collected with a gas-tight syringe and analyzed with a gas chromatograph with a flame ionization detector and a fused silica column. Samples were periodically obtained for analysis by gas chromatography/ mass spectroscopy.

Exposures were stagger-started over a 5-day period with 16 animals, 2/sex/group, receiving their first exposure on each of 5 consecutive days.

The rats were observed for signs of toxicity daily prior to initiation of

5. Toxicity Id 108-20-3

Date

exposures, and individual body weights were recorded weekly.

During weeks of Functional Observation Battery (FOB) evaluation, four rats/group would be removed from the inhalation chamber and housed separately overnight and evaluated on the following day between the hours of 7:30 a.m. and 11:00 a.m. The process was repeated for each consecutive day until all rats were evaluated. The rats were evaluated with minimal disruption to the exposure schedule and with a manageable number of rats per day, 16 on any given day. The animals were evaluated in an FOB followed by a determination of motor activity prior to initiation of exposure; the FOB following 2, 4, 8, and 13 weeks of exposures, and for motor activity following 4, 8, and 13 weeks of exposures. Following the final determination of motor activity, the animals were anesthetized, intravascularly perfused, and the brain, spinal cord, and peripheral nerves removed for microscopic examination.

The FOB consisted of initially observing home-cage positioning, posture, and reaction to removal from the cage. This was followed by evaluation for exophthalmus/palpebral closure, lacrimation, salivation, pupillary response, palpebral reflex, and pinna reflex. These observations were scored by type and intensity. The animals were then observed for open field behavior. Piloerection, respiratory rate, tremors, convulsions, posture, gait, ataxic gait, tail elevation, unperturbed activity level, vocalization, number of rears, fecal balls, and urine pools were all recorded during the open-field observations. Reactions to the approach of a pencil, finger snap, and tail pinch were ranked and recorded. Finally, fore- and hindlimb grip strength, rectal temperature, and body weight were measured. Automated motor activity was assessed for 30 minutes in figure-eight mazes after the completion of the FOB.

Following the last FOB and motor activity evaluation, the rats were anesthetized with heparinized sodium pentobarbital (i.p.). The thoracic cavity was opened and the animals were infused with phosphate-buffered gluteraldehyde through the left ventricle. The perfused brain, spinal cord, and sciatic nerve with its tibial, sural, and peroneal divisions were removed. The brain and nerve tissues were processed for embedding in paraffin or glycol methacrylate (dorsal root ganglia and peripheral nerves) and sectioned for light or electron microscopic pathologic evaluation.

Type: 90-Day Neurotoxicity

Species/strain: Rat/Sprague-Dawley; Tac:N(SD)fBR

No./sex/dose: 10/sex/group

Vehicle: None

Method: USEPA 1984, 40CFR Part 798:6050, 6400, and 6200

Motor activity in a figure-eight maze and unperturbed activity in the FOB were decreased at week 4 in females exposed to 7060 ppm; activity in the FOB was also decreased in females exposed to 450 ppm at week 4. Other changes in the FOB appeared to be minor, and no changes were observed

during microscopic examination of tissues from the nervous system.

: Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.

: Inhalation exposures to DIPE at concentrations as high as 7060 ppm for 13

weeks resulted in few observable effects on the nervous system.

Reliability : (2) valid with restrictions

GLP unknown. Study well documented, meets generally accepted scientific

principles, acceptable for assessment.

01.11.2005 (35)

5.5 GENETIC TOXICITY 'IN VITRO'

Result

Test substance

Conclusion

Type : Bacterial reverse mutation assay

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System of testing : Salmonella typhimurium

Test concentration: Up to 8000 ug/ml in the pre-incubation mix

Cycotoxic concentr. :

Metabolic activation: with and without

Result : negative

Method : other: Similar to OECD Guideline 471

Year : 1988 GLP : no data

Test substance : other TS: Diisopropyl ether (CAS No. 108-20-3)

Remark: Strains tested: Salmonella typhimurium tester strains TA98, TA100,

TA1535, TA1537, TA1538

Exposure method: Preincubation assay for volatile compounds [Brooks

and Dean 1981]

Test Substance Doses/concentration levels: Up to 8000 ug/ml in the pre-

incubation mix

Metabolic activation: With and without (S9 fraction mix of livers of Aroclor

1254 pretreated rats)

Vehicle: Tween 80/ethanol

Tester strain, activation status, Positive Controls and concentration level:

Not stated

Statistical analysis: Mean revertant colony count and standard deviation

were determined for each dose point.

Dose Rangefinding Study: Cytotoxicity study

S9 Optimization Study: No

Result: DIPE did not induce reverse gene mutation in any strain. The test

substance was not genotoxic in this assay with or without metabolic

activation.

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis

[propane], 2,2 '-

Source/purity not specified.

Conclusion: Under the conditions of this study, the test material was not mutagenic.

Reliability : (2) valid with restrictions

Restriction due to the lack of any information regarding the selection of dose levels used during the study. In addition no information is presented regarding cytotoxicity or the presence of test material precipitate in the cultures. Although study was not stated to be conducted under GLP, it was conducted at a reputable laboratory [Shell Research Limited, Sittingbourne

Research Center].

10.11.2005 (5)

Type : Sister chromatid exchange assay System of testing : Chinese hamster ovary cells

Test concentration : Up to 1200 ug/ml

Cycotoxic concentr.

Metabolic activation: withoutResult: negative

Method : other: Similar to OECD Guideline 473

Year : 1984 GLP : no data

Test substance : other TS: Di-isopropyl ether (CAS No. 108-20-3)

Remark : Test type: Chromosome damage

5. Toxicity Id 108-20-3

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Exposure method: For volatile compounds

Metabolic activation: Metabolic activation S9 was not added because liver

cells are metabolically competent

Vehicle: Tween 80/ethanol

Tester strain, activation status, Positive Controls and concentration level: Cultured CHO cells were grown in 80 cm2 flasks for 24 hr before compound treatment. Treatment periods were 5 hr in the presence of S9 mix and 24 hr in the absence of S9. Colcemid was added to all cultures 22 hr after the initial treatment. After a further 2 hr, the cells were trypsinized, resuspended in hypotonic solution and then fixed, before spotting onto slides. Cell preparations were then stained with Giemsa. The slides were randomly coded and 100 cells from each culture were analyzed microscopically. Mitotic index estimations were also made. The positive controls were ethylmethanesulfonate [-S9] and cyclophosphamide [+S9].

Vehicle control: Yes

Dose rangefinding study: Cytotoxicity study

S9 Optimization study: No

Result: DIPE did not induce chromosomal damage in CHO cells. The test

substance was not genotoxic in this assay.

Test substance : Di-isopropyl ether (CAS No. 108-20-3)

Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis

[propane], 2,2 '-

Source/purity of test material: 98.5%

Conclusion : Under the conditions of this study, the test material was not mutagenic.

Reliability : (2) valid with restrictions

Restriction due to the lack of any information regarding the selection of dose levels used during the study. In addition no information is presented regarding cytotoxicity or the presence of test material precipitate in the cultures. Although study was not stated to be conducted under GLP, it was conducted at a reputable laboratory [Shell Research Limited, Sittingbourne

Research Center].

10.11.2005 (4)

Type : DNA damage and repair assay

System of testing : Rat liver cells
Test concentration : Up to 1200 ug/ml

Cycotoxic concentr. :

Metabolic activation: withoutResult: negative

Method : other: Similar to OECD Guideline 476

Year : 1984 GLP : no data

Test substance: other TS: Di-isopropyl ether (CAS No. 108-20-3)

Remark: Test type: Chromosome damage

Strains tested: RL4

Metabolic activation: Metabolic activation S9 was not added because liver

cells are metabolically competent.

Vehicle: Tween 80/ethanol

Tester strain, activation status, Positive Controls and concentration level: Cultured rat liver cells were grown and treated on glass microscope slides 5. Toxicity Id 108-20-3

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contained in 100 ml glass Leighton tubes. After 22 hr exposure to test compound or solvent, colcemid was added to each culture. After a further 2 hr, the slides were removed, subjected to hypotonic treatment followed by fixation and stained with Giemsa. The preparations were randomly coded and 100 cells from each culture were analyzed microscopically. The positive control was 7,12-dimethylbenzanthracenene.

Vehicle control: Yes

Dose rangefinding study: Cytotoxicity study

S9 Optimization study: None needed

Result: DIPE did not induce chromosomal damage in rat liver cells. The test

substance was not genotoxic in this assay.

Test substance: Di-isopropyl ether (CAS No. 108-20-3)

Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis [propane],

2,2 '-

Source/purity of test material: 98.5%

Conclusion : Under the conditions of this study, the test material was not mutagenic.

Reliability : (2) valid with restrictions

Restriction due to the lack of any information regarding the selection of dose levels used during the study. In addition no information is presented regarding cytotoxicity or the presence of test material precipitate in the cultures. Although study was not stated to be conducted under GLP, it was conducted at a reputable laboratory [Shell Research Limited, Sittingbourne

Research Center].

10.11.2005 (4)

Type : Gene mutation in Saccharomyces cerevisiae

System of testing : Saccharomyces cerevisiae

Test concentration: Up to 8000 ug/ml in the pre-incubation mix

Cycotoxic concentr.

Metabolic activation: with and without

Result : negative

Method : other: Similar to OECD Guideline 481

Year : 1984 GLP : no data

Test substance: other TS: Di-isopropyl ether (CAS No. 108-20-3)

Remark: Test type: Yeast mitotic gene conversion

Strains tested: JD1

Exposure method: [Brooks and Dean 1981]

Metabolic activation: With and without (S9 fraction mix of livers of Aroclor

1254 pretreated rats)

Vehicle: Tween 80/ethanol

Tester strain, activation status, Positive Controls and concentration level: Yeast cells were grown in log-phase, washed and resuspended in 2/5 strength YEPD broth at a concentration of 1 X 107 cells/ml. The suspension was divided into 1.9 ml amounts in 30 ml universal containers and 0.1 ml of test compound solution was added. For experiments with metabolic activation [+S9], 0.1 ml of DIPE was added to 01.6 ml of yeast cell suspension, together with 0.3 ml of S9 mix. Initially a range of concentrations of DIPE was tested up to 5 mg/ml if solubility allowed. A second experiment was performed based on these results and taking into account cell viability. The cultures were incubated with shaking at 30 C for 18 hr. Aliquots were plated onto the appropriate culture media for selection of mitotic gene convertants and cells surviving the treatment. Mitotic gene

5. Toxicity Id 108-20-3

Date

conversion may be scored by supplementing the minimal medium with histidine to score tryptophan prototrophs, and with tryptophan to score histidine prototrophs. Control plates were set up with solvent alone and with the positive control compounds 4-nitroquinoline oxide and

cyclophosphamide.

Vehicle control: Yes

Dose rangefinding study: Cytotoxicity study

S9 Optimization study: No

Result: DIPE did not induce mitotic gene conversion I yeast. The test substance

was not genotoxic in this assay with or without metabolic activation.

Test substance: Di-isopropyl ether (CAS No. 108-20-3)

Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis [propane],

2.2 '-

Source/purity of test material: 98.5%

Conclusion: Under the conditions of this study, the test material was not genotoxic.

Reliability : (2) valid with restrictions

Restriction due to the lack of any information regarding the selection of dose levels used during the study. In addition no information is presented regarding cytotoxicity or the presence of test material precipitate in the cultures. Although study was not stated to be conducted under GLP, it was conducted at a reputable laboratory [Shell Research Limited, Sittingbourne

Research Center].

10.11.2005 (4)

5.6 GENETIC TOXICITY 'IN VIVO'

5.7 CARCINOGENICITY

5.8.1 TOXICITY TO FERTILITY

5.8.2 DEVELOPMENTAL TOXICITY/TERATOGENICITY

Species : rat Sex : female

Strain : Sprague-Dawley
Route of admin. : inhalation
Exposure period : 6 hr/day

Frequency of treatm. : Gestation Days 6-15

Duration of test : 20 days

Doses : 0, 430, 3095, or 6745 ppm

Control group : other: yes (untreated & sham-exposed)

other: NOEL Maternal : = 430 ppm other: NOEL Pup : = 430 - ppm

Result : Maternal NOEL: 430 ppm; Pup NOEL: 430 ppm

Method : EPA OTS 798.4350

Year : 1996 GLP : no data

Test substance : other TS: Diisopropyl Ether (CAS No. 108-20-3)

Method : Statistical method:

Statistical analyses of numerical data included ANOVA and Tukey's studentized range test for data on serum chemistry. Duncan's multiple

Id 108-20-3 5. Toxicity Date 27.02.2006

> range test was used for hematology and body weights to assess statistically significant differences between control and exposed groups. Data on the maternal biophase, cesarean sections, and fetuses were evaluated by ANOVA followed by group comparisons using Fisher's exact or Dunnett's test.

Remark

Nulliparous females were housed with males in a 1:1 ratio and observed daily for evidence of breeding activity. Females positive for sperm plug and for sperm in the vaginal lavage fluid were considered to be at day 0 of gestation and were individually housed. The females were then randomly distributed to 5 groups of 22 animals each; untreated controls, shamexposed controls, and 3 groups exposed to vapors of DIDP at 430, 3095, or 6745 ppm for 6 hr/day on gestation days (GD) 6-15.

Vapors were generated by metering DIPE on to a warmed fiberglass wick and carried to the three 1m3 exposure chambers by a stream of measured and filtered room air. Temperature and humidity in the chambers were measured every 30 minutes and air flow through each chamber was metered to provide at least 12 air changes/hour. Chamber air samples were collected with a gas-tight syringe and analyzed with a gas chromatograph with a flame ionization detector and a fused silica column. Samples were periodically obtained for analysis by gas chromatography/ mass spectroscopy.

Sham-exposed and test animals were housed in their exposure chambers throughout the exposure period; untreated controls in a separate room. Food and water were not available during the 6 -hour exposure periods but were available ad libitum at all other times. All animals were observed daily. Body weights were recorded on days 0, 6, 13, 16, and 20. Food consumption was measured on GD 6, 13, 16, and 20. Females were sacrificed on GD 20 by diethyl ether overexposure followed by exsanguination. Serum samples from the descending aorta were analyzed for glucose, urea nitrogen, total protein, albumin, globulin, A/G ratio, sorbitol dehydrogenase, lactate dehydrogenase, aspartate aminotransferase, alanine aminotransferase, alkaline phosphatase, total bilirubin, creatinine, cholesterol, triglycerides, uric acid, Cl, Ca, Na, K, and P. All organs were examined grossly. The number of corpora lutea per ovary and the gravid uterine weights were recorded. Uterine contents were examined and the numbers of implantation sites, early and late resorptions, and live and dead fetuses were recorded. The gender of each fetus was recorded. Fetuses were weighed and examined for gross anomalies. Fetuses of each litter were equally distributed between two groups; half were fixed in Bouin's solution and examined for visceral anomalies and the remaining fetuses were fixed in 95% ethanol and examined for skeletal anomalies after differential staining for cartilage and bone.

Dams exposed to 6745 ppm had a slight reduction in body weight gain and a significant decrease in food consumption. A concentration-related increase in the incidence of rudimentary 14th ribs was observed (statistically significant at 3095 and 6745 ppm) but the relevance of the finding was uncertain. There was no apparent toxicity, either maternal or fetal, at the lowest exposure concentration, 430 ppm.

Type: Developmental Toxicity

Species/strain: Sprague-Dawley; VAF/Plus Crl:CD(SD)BR

No./dose: 22/group Vehicle: None

Method: USEPA 1984; 40CFR Part 798:4350

Maternal NOEL: 430 ppm Result Pup NOEL: 430 ppm

Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.

Test substance Conclusion DIPE is not a teratogen. Reliability (2) valid with restrictions

5. Toxicity Id 108-20-3

Date

GLP unknown. Study well documented, meets generally accepted scientific

principles, acceptable for assessment.

01.11.2005

5.8.3 TOXICITY TO REPRODUCTION, OTHER STUDIES

5.9 SPECIFIC INVESTIGATIONS

5.10 EXPOSURE EXPERIENCE

5.11 ADDITIONAL REMARKS

Type: other: Sensory Irritation in Humans

Method : Non-guideline.

Remark : Species/strain: Humans Sex: Male and female

Number/sex/group: Average of 12

Route of administration: Inhalation

Vehicle: None Control: No Year: Prior to 1946

GLP: No

Result: 300 ppm: 35% of the subjects objected to this solvent because of the

unpleasant odor rather than irritation.

500 ppm: there was a sensory response that was acceptable to the

majority of subjects.

Test condition: Subjects were exposed for 15 minutes and olfactory fatigue and irritation of

mucous membranes were reported. "Motion pictures were shown to occupy the subject's attention and divert their thoughts from the atmospheric

contamination to which they were exposed."

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis

[propane], 2,2 '-

Test material is stated to be technical grade product.

Conclusion : DIPE does not appear to be a sensory irritant at concentrations up to 500

ppm, but it does have an unpleasant odor at this concentration.

Reliability : (2) valid with restrictions

Not GLP but conducted at a reputable laboratory [Harvard School of Public

Health, Boston].

01.11.2005 (37)

Type : other: Sensory irritation in humans

Method : Non-Guideline.

Remark : Species/strain: Young adult humans [University of California staff and

medical students]
Sex: Not specified

Number/sex/group: Not specified Route of administration: Inhalation

Vehicle: None Control: No Year: 1955 GLP: No

Result : Numbers of subjects with degree of effect

5. Toxicity Id 108-20-3
Date 27.02.2006

Concentration 400 ppm 800 ppm

Number subjects: 7 7

Eye irritation: 7 absent 3 absent, 3 slight, 1 mod. Nose irritation: 5 absent, 2 slight 2 absent, 5 slight Pulmonary discomfort: 7 absent 4 absent, 3 slight Olfactory cognition: 1 slight, 6 mod. 4 mod., 3 severe

CNS effects: 7 absent 7 absent

Test condition : Exposures were conducted in a whole-body chamber approximately 7700 l

equipped with a fan. Exposures were made in a static atmosphere

generated by vaporizing a predetermined quantity of test solvent from a hot surface. Five minutes were allowed for evaporation and equilibration, and subjects were exposed for 5 minutes, during which time they noted the

degree of subjective responses at one-minute intervals.

Test substance: Diisopropyl ether (CAS No. 108-20-3)

Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis

[propane], 2,2 '-

Test material is stated to be commercial grade with purity of 98% or better,

provided by Shell Chemical Corporation.

Conclusion : 400 ppm: 5 mins of inhalation exposure caused no eye irritation, none to

slight nose irritation, no pulmonary discomfort, olfactory recognition but no

central nervous system effects.

800 ppm: 5 mins of inhalation exposed caused slight eye and nose

irritation, none to slight pulmonary discomfort, definite olfactory recognition

but no central nervous system effects.

Reliability : (2) valid with restrictions

Not GLP but conducted at a reputable laboratory [University of California

School of Medicine].

01.11.2005 (22)

6. Analyt. Meth. for Detection and Identification	ld 108-20-3 Date
6.1 ANALYTICAL METHODS	
6.2 DETECTION AND IDENTIFICATION	
49 / 55	

7. Eff	. Against Target Org. and Int	ended Uses	108-20-3 27.02.2006	
7.1	FUNCTION			
7.2	EFFECTS ON ORGANISMS TO BE CON	TROLLED		
7.3	ORGANISMS TO BE PROTECTED			
7.4	USER			
7.5	RESISTANCE			
		50 / 55		

Id 108-20-3 8. Meas. Nec. to Prot. Man, Animals, Environment Date 27.02.2006 8.1 METHODS HANDLING AND STORING 8.2 FIRE GUIDANCE **EMERGENCY MEASURES** 8.3 POSSIB. OF RENDERING SUBST. HARMLESS 8.4 **WASTE MANAGEMENT SIDE-EFFECTS DETECTION** 8.6 8.7 SUBSTANCE REGISTERED AS DANGEROUS FOR GROUND WATER 8.8 REACTIVITY TOWARDS CONTAINER MATERIAL

9. References Id 108-20-3

Date

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(46)

10. Summary and Evaluation	ld 108-20-3
	Date 27.02.2006
10.1 END POINT SUMMARY	
40.0 HAZADD OHUMADY	
10.2 HAZARD SUMMARY	
10.3 RISK ASSESSMENT	

RECEIVED OPPT CBIC

2007 JAN -4 AM 7: 34

201-16472D

IUCLID

Data Set

 Existing Chemical
 : ID: 78-92-2

 CAS No.
 : 78-92-2

 EINECS Name
 : butan-2-ol

 EC No.
 : 201-158-5

 TSCA Name
 : 2-Butanol

Producer related part

Molecular Formula

Company : ExxonMobil Biomedical Sciences Inc.

: C4H10O

Creation date : 09.01.2002

Substance related part

Company : ExxonMobil Biomedical Sciences Inc.

Creation date : 09.01.2002

Status Memo

Printing date : 01.06.2006

Revision date :

Date of last update : 01.06.2006

Number of pages : 76

Chapter (profile) : Chapter: 1, 2, 3, 4, 5, 6, 7, 8, 10
Reliability (profile) : Reliability: without reliability, 1, 2, 3, 4

Flags (profile) : Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE),

Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

Id 78-92-2

Date

1.0.1 APPLICANT AND COMPANY INFORMATION

Type :

Name : Atochem

Contact person

Date

Street : 4, Cours Michelet
Town : 92080 Paris la Defense

Country : France

Phone

Telefax :
Telex :
Cedex :
Email :
Homepage :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type :

Name : Deutsche Shell Chemie GmbH

Contact person

Date

Street : Koelner Strasse 6
Town : 65760 Eschborn

Country : Germany

Phone

Telefax
Telex
Cedex
Email
Homepage

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Туре

Name : EXXON CHEMICAL, Limited

Contact person

Date

Street : 4600 Parkway

Town: PO15 7AP Fareham, Hampshire

 Country
 : United Kingdom

 Phone
 : (44)489.88.4480

 Telefax
 : (44)489.88.4455

Telex :
Cedex :
Email :
Homepage :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type :

Name : RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Contact person

Date

Street : Ueberseering 40
Town : 22297 Hamburg

Id 78-92-2

Date

 Country
 : Germany

 Phone
 : 040-6375-0

 Telefax
 : 040-6375-3496

 Telex
 : 21151320

Cedex : Email : Homepage :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type :

Name : SHELL FRANCE

Contact person

Date

Street : 89 bld Franklin Roosevelt
Town : 92564 Rueil Malmaison

Country : France

Phone : 33 1 47.14.71.00 Telefax : 33 1 47.14.82.99 Telex : SHELL 615013F

Cedex : Email : Homepage :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type

Name : Shell Nederland Chemie B.V.

Contact person

Date

Street : P.O. Box 3030

Town : 3190 GH Hoogvliet-Rotterdam

Country : Netherlands
Phone : +31-10-2317005
Telefax : +31-10-2317125

Telex : Cedex : Email : Homepage :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type

Name : Union Carbide Benelux

Contact person

Date :

Street : Norderlaan 147
Town : 2030 Antwerpen

Country : Belgium

Phone
Telefax
Telex
Cedex
Email

Homepage :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Id 78-92-2

Date

1.0.2 LOCATION OF PRODUCTION SITE, IMPORTER OR FORMULATOR

1.0.3 IDENTITY OF RECIPIENTS

1.0.4 DETAILS ON CATEGORY/TEMPLATE

1.1.0 SUBSTANCE IDENTIFICATION

1.1.1 GENERAL SUBSTANCE INFORMATION

Purity type :

Substance type : organic Physical status : liquid

Purity
Colour
Odour

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

1.1.2 SPECTRA

1.2 SYNONYMS AND TRADENAMES

2-Butanol

Source : Shell Nederland Chemie B.V. Hoogvliet-Rotterdam

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

19.05.1994

2-butanol

Source : Union Carbide Benelux Antwerpen

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

26.05.1994

2-Hydroxybutane

Source : Shell Nederland Chemie B.V. Hoogvliet-Rotterdam

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

19.05.1994

butan-2-ol

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Hamburg

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

16.02.1994

Ethyl methyl carbinol

Id 78-92-2

Date

Source : Shell Nederland Chemie B.V. Hoogvliet-Rotterdam

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

19.05.1994

SBA

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Hamburg

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

16.02.1994

SBA; sec-Butanol; Butan-2-ol; secondary butyl alcohol

Source : Atochem Paris la Defense

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

27.04.1994

SBA; secondary butyl alcohol; ethyl methyl carbinol; 2-hydroxybutane

Source : SHELL FRANCE Rueil Malmaison

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

20.05.1994

sec-butanol

Source : Union Carbide Benelux Antwerpen

EXXON CHEMICAL, Limited Fareham, Hampshire RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Hamburg

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

26.05.1994

sec.-butyl alcohol, iso-butanol

Source : Deutsche Shell Chemie GmbH Eschborn

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

30.05.1994

secondary butyl alcohol

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Hamburg

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

16.02.1994

secundary butyl alcohol

Source : Union Carbide Benelux Antwerpen

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

26.05.1994

1.3 IMPURITIES

1.4 ADDITIVES

Id 78-92-2

Date

1.5 TOTAL QUANTITY

Quantity : 100000 - 500000 tonnes in

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

1.6.1 LABELLING

Labelling : as in Directive 67/548/EEC

R-Phrases : (10) Flammable

(36/37) Irritating to eyes and respiratory system (67) Vapours may cause drowsiness and dizziness

S-Phrases : (2) Keep out of reach of children

(7/9) Keep container tightly closed and in a well-ventilated place (13) Keep away from food, drink and animal feeding stuffs

(24/25) Avoid contact with skin and eyes

(26) In case of contact with eyes, rinse immediately with plenty of water

and seek medical advice

(46) If swallowed, seek medical advice immediately and show this

container or label

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

1.6.2 CLASSIFICATION

Classified: as in Directive 67/548/EEC

Class of danger : irritating

R-Phrases : (36/37) Irritating to eyes and respiratory system

Specific limits :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Classified : as in Directive 67/548/EEC

Class of danger

R-Phrases : (10) Flammable

Specific limits

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Classified: as in Directive 67/548/EEC

Class of danger

R-Phrases: (67) Vapours may cause drowsiness and dizziness

Specific limits :

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Id 78-92-2

Date

1.6.3 PACKAGING

1.7 USE PATTERN

Type of use : type

Category : Non dispersive use

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : type

Category : Use in closed system

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : type

Category : Use resulting in inclusion into or onto matrix

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : type

Category : Wide dispersive use

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : industrial

Category : Basic industry: basic chemicals

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : industrial

Category: Chemical industry: used in synthesis

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : industrial Category : Fuel industry

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : industrial

Category: Paints, lacquers and varnishes industry

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : use

Category : Intermediates

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : use

Id 78-92-2

Date

Category : Solvents

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : use

Category : other: fuel additive in lead-free gasoline

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

Type of use : use Category : other

Source : EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

11.02.2000

1.7.1 DETAILED USE PATTERN

1.7.2 METHODS OF MANUFACTURE

1.8 REGULATORY MEASURES

1.8.1 OCCUPATIONAL EXPOSURE LIMIT VALUES

Type of limit : MAC (NL) Limit value : 450 mg/m3

Source : Shell Nederland Chemie B.V. Hoogvliet-Rotterdam

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

03.06.1994 (29

Type of limit : MAC (NL)
Limit value : 450 mg/m3

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

16.02.1994

Type of limit : MAC (NL) Limit value : 450 mg/m3

Source : RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Hamburg

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

13.05.1994

Type of limit : MAK (DE) Limit value : 300 mg/m3

Short term exposure limit value

Limit value: 600 mg/m3Time schedule: 30 minute(s)Frequency: 4 times

Source : Atochem Paris la Defense

ld 78-92-2

Date

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

27.04.1994

MAK (DE) Type of limit Limit value 300 mg/m3

Source EXXON CHEMICAL, Limited Fareham, Hampshire

RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Hamburg

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

16.02.1994

Type of limit MAK (DE) Limit value 100 ml/m3

Remark Substance is easily resorbed.

Deutsche Shell Chemie GmbH Eschborn Source

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

30.05.1994

Type of limit : OES (UK) Limit value 300 mg/m3

Short term exposure limit value

Limit value 450 mg/m3

Time schedule

Frequency times

Source EXXON CHEMICAL, Limited Fareham, Hampshire

RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

16.02.1994

Type of limit TLV (US) Limit value 303 mg/m3

Shell Nederland Chemie B.V. Hoogvliet-Rotterdam Source

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

03.06.1994 (3)

Type of limit TLV (US) Limit value 303 mg/m3

Short term exposure limit value

Limit value : 455 mg/m3

Time schedule

Frequency times

Source : Union Carbide Benelux Antwerpen

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

26.05.1994 (1)

Type of limit TLV (US) Limit value 303 mg/m3

Short term exposure limit value

: 455 mg/m3 Limit value 15 minute(s) Time schedule Frequency 4 times

Source Atochem Paris la Defense

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

27.04.1994 (53)

Id 78-92-2

Date

Type of limit : TLV (US) Limit value : 303 mg/m3

Remark: Use local exhaust ventilation.

Hand protection: PVC, nitrile or neoprene gloves.

Eye protection: safety monogoggles.

Body protection: standard issue work clothes.

chemicals resistant safety shoes or boots.

Source : SHELL FRANCE Rueil Malmaison

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

26.05.1994 (88)

Type of limit : TLV (US) Limit value : 303 mg/m3

Short term exposure limit value

Limit value : 455 mg/m3 Time schedule : 8 hour(s) Frequency : times

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

RWE-DEA Aktiengesellschaft für Mineraloel und Chemie

Hamburg

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

16.02.1994

Type of limit : other: VME

Limit value : 300

Country : France

Source : Atochem Paris la Defense

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

27.04.1994 (52)

1.8.2 ACCEPTABLE RESIDUES LEVELS

1.8.3 WATER POLLUTION

1.8.4 MAJOR ACCIDENT HAZARDS

1.8.5 AIR POLLUTION

1.8.6 LISTINGS E.G. CHEMICAL INVENTORIES

1.9.1 DEGRADATION/TRANSFORMATION PRODUCTS

1.9.2 COMPONENTS

1.10 SOURCE OF EXPOSURE

Id 78-92-2

Date

Remark

: As the quantities of this substance placed on the EU market by Union Carbide Benelux N.V. are normally sourced from the manufacturing facilities of its U.S. parent company, no exposure can arise within the EU from the manufacture of these substances. The comments below on exposure are restricted to uses for which Union Carbide believes its customers use this substance.

Major use(s): Fuel additive in lead-free gasoline.

Sources of human exposure: Very minor sporadic exposure to

the public via inhalation during filling of vehicles.

Quantitative estimates are not available.

Sources of environmental exposure: Negligible sporadic exposure to the atmosphere during filling of vehicles. Quantitative estimates are not available. Substance is essentially oxidised to carbon dioxide and water during

use.

Source : Union Carbide Benelux Antwerpen

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

24.05.1994

Remark: Continuous process. Non-direct hydratation by absorption of

butenes in sulfuric acid. One production site.

Distribution pattern: from production

% to air 0.02 % to water 0.02 to soil 0.0 to sediment 0.0

Source : Atochem Paris la Defense

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

03.06.1994

Remark: Inhalation or skin contact when loading, unloading, using

the product.

In case of accidental release, product may contaminate the

environment.

Source : SHELL FRANCE Rueil Malmaison

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

23.02.1994

1.11 ADDITIONAL REMARKS

Remark : DISPOSAL

Recover or recycle if possible. Otherwise incineration.

TRANSPORT INFORMATION

UN Number: 1120

Class: 3

Packing Group: III

Proper Shipping Name: Secondary butyl alcohol

Date

Sea (IMO) Class: 3.3

Packing Group: III

Symbol: Flammable liquid Marine Pollutant (Y/N): No

Rail/Road (RID/ADR)

Class: 3 Item: 31(c)

Symbol: Flammable liquid Kemler Plate: 30/1120

Air (IATA/ICAO)

Class: 3

Packing Group: III

Symbol: Flammable liquid

Source : Shell Nederland Chemie B.V. Hoogvliet-Rotterdam

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

10.06.1994

Remark: Disposal: Incinerate in a furnace where permitted under

national and local regulations.

Transport: Butan-2-ol is classified as a class 3 product according the ADR/RID/IMDG/ICAO regulations. Butan-2-ol is shipped in road/rail tankcars.

tankcontainers/ISOtanks and smaller packages (e.g. drums).

ż.

Source : Union Carbide Benelux Antwerpen

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

26.05.1994

Remark: Recover or recycle if possible. Otherwise: incineration.

Avoid electrostatic discharge generation.

Earth all equipment. Avoid splash filling.

Use a vapour recovery system. Keep in a well ventilated place.

Avoid naked flames. Remove ignition sources.

Do not smoke. Avoid sparks.

Transport

UN number: 1120

Class: 3

Packing Group: III

Proper Shipping Name: Secondary butyl alcohol

Sea (IMO) Class: 3.3

Marine Pollutant : No Symbol : Flammable liquid

Rail/Road (ADR/RID)

Class: 3 Item: 31 c)

Symbol : Flammable liquid Kemler Plate : 30/1120

Air (IATA/IACO)

Id 78-92-2

Date

Class: 3

Symbol: Flammable liquid

Source : SHELL FRANCE Rueil Malmaison

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

26.05.1994

Remark: Disposal options:

Recover or recycle if possible. Otherwise: incineration.

Transport classification:

UN number: 1120

ADR/RID:

class/item: 3/31 c packing group: 3 Kemler number: 30

label: 3

Proper shipping name: sec.-butyl alcohol

ICAO: class: 3

packing group: III label: flammable liquid

Proper shipping name: butanols

IMO/IMDG: class: 3.3

packing group: III Marine pollutant: no label: flammable liquid IMDG page: 3313 EMS number: 3-06 MFAG plate: 305

Source : Deutsche Shell Chemie GmbH Eschborn

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

30.05.1994

1.12 LAST LITERATURE SEARCH

1.13 REVIEWS

Id 78-92-2

Date

2.1 MELTING POINT

Value : = -114 °C

Sublimation

Method : other: no data

Year :

GLP : no data

Test substance

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

17.01.2002 (50)

Value : ca. -89 - -108 °C

Sublimation

Method : other: no data

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (57)

2.2 BOILING POINT

Value : = 99.5 °C at

Decomposition

Method : other: no data

Year :

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

17.01.2002 (38)

Value : ca. 99 - 102 °C at 1013 hPa

Decomposition

Method : other: ASTM D1078/86

Year : 1986 GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002

Id 78-92-2

Date

2.3 DENSITY

Type : density

Value : = 806 kg/m 3 at 20 °C

Method : other: no data

Year :

GLP : no data

Test substance

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonBiomedical Sciences, Inc., New Jersey, USA

17.01.2002 (23)

Type : density

Value : ca. 807 kg/m3 at 15 °C

Method : other: ASTM D4052/86

Year : 1986 GLP : no data

Test substance :

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (39)

Type : density

Value : = 3.29 kg/m3 at 20 °C

Method : other: no data

Year :

GLP : no data

Test substance :

Remark: Vapour density

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (76)

2.3.1 GRANULOMETRY

2.4 VAPOUR PRESSURE

Value : = 15.98 hPa at 20 °C

Decomposition

Method : other (measured)

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (57)

Id 78-92-2

Date

2.5 **PARTITION COEFFICIENT**

Partition coefficient

Log pow = .61 at 20 °C

pH value

Method

Year

other (measured)

no data

GLP

Test substance

Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source

Remark

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

17.01.2002

(47)

2.6.1 SOLUBILITY IN DIFFERENT MEDIA

Solubility in

Value

pH value

= 125 g/l at 20 °C

concentration at °C

Temperature effects

Examine different pol.

pKa at 25 °C

Description

Stable

Deg. product

Method other: no data

Year

GLP no data

Test substance

: Information on sBA purity is not available; assumed to have used a Remark

commercial grade of >/= 99%.

EXXON CHEMICAL, Limited Fareham, Hampshire Source

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

(46)17.01.2002

Solubility in

Value = 181 g/l at 25 °C

pH value

concentration at °C

Temperature effects

Examine different pol.

pKa at 25 °C

Description

Stable

Deg. product

Method other: no data

Year

GLP no data

Test substance

Remark : Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA Source

17.01.2002 (51)

Id 78-92-2

Date

Solubility in

Value ca. 201 g/l at 20 °C

pH value

at °C concentration

Temperature effects

Examine different pol.

pKa at 25 °C

Description **Stable**

Deg. product

Method

other: no data

Year

GLP no data

Test substance

Information on sBA purity is not available; assumed to have used a Remark

commercial grade of >/= 99%.

Source EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (57)

2.6.2 SURFACE TENSION

2.7 FLASH POINT

= 24 °C Value Type closed cup

other: NFT 60 103 Method

Year

GLP no data

Test substance

Remark : Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (74)

Value : ca. 25 °C Type closed cup

Directive 84/449/EEC, A.9 "Flash point" Method

Year 1987 **GLP** no data

Test substance

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

01.06.1994

AUTO FLAMMABILITY 2.8

: > 350 °C at 1013 hPa Value

Method : other: E695/85

Year 1985 **GLP** : no data

Id 78-92-2

Date

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (39)

Value : = 406 °C at 1013 hPa

Method : other: no data

Year :

GLP : no data

Test substance :

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

02.06.1994 (75)

2.9 FLAMMABILITY

Result : flammable

Method : other: no data

Year :

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (74)

2.10 EXPLOSIVE PROPERTIES

Method : other: no data

Year :

GLP : no data

Test substance :

Remark: Explosive limits of vapours in air: 1.7 to 9.8% vol.

Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (74)

2.11 OXIDIZING PROPERTIES

2.12 DISSOCIATION CONSTANT

2.13 VISCOSITY

2. Physico-Chemical Data	ld Date	78-92-2
2.14 ADDITIONAL REMARKS		
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3. Environmental Fate and Pathways

Date

3.1.1 PHOTODEGRADATION

Type : air Light source :

Light spectrum: nm

Relative intensity : based on intensity of sunlight

INDIRECT PHOTOLYSIS

Sensitizer : OH

Conc. of sensitizer : 1500000 molecule/cm³

Rate constant : = $.000000000099751 \text{ cm}^3/(\text{molecule*sec})$

Degradation : = 50 % after 12.9 hour(s)

Deg. product

Method : other (calculated)

Year : GLP : Test substance :

Remark : 50% degradation after 1.07 days based on a 12-hr. day. The SMILES

(Simplified Molecular Input Line Entry System) structure used with the

model was: OC(CC)C.

Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Reliability : (2) valid with restrictions

Rated a 2 for reliability because it is a calculated value.

17.01.2002 (37)

Type : air Light source :

Light spectrum : nm

Relative intensity : based on intensity of sunlight

INDIRECT PHOTOLYSIS

Sensitizer : OH

Conc. of sensitizer

Rate constant : cm³/(molecule*sec)

Degradation: % after

Deg. product

Method : other (measured)

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Rate constants were determined from data developed in three labs. Two equations were applied to the measured data and designated the TI (time-included) and TE (time-excluded) methods. Both equations are in the form of a straight line equation, y=mx+b. Least square linear regression

analyses were applied to the data from each experiment.

Result : Hydroxyl rate constants as calculated from measured data using the time-

included equation:

 $10.30 + -2.48 \times 10-12 \text{ cm}3/\text{molecule-sec (r2} = 0.97); @ 24C 11.55 + -1.77 \times 10-12 \text{ cm}3/\text{molecule-sec (r2} = 0.99); @ 24C 4.01 + -1.38 \times 10-12 \text{ cm}3/\text{molecule-sec (r2} = 0.76); @ 34C$

Hydroxyl rate constants as calculated from measured data using the time-

excluded equation:

3. Environmental Fate and Pathways

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Date

9.40 +/- 0.87 x 10-12 cm3/molecule-sec (r2 = 0.99); @ 24C 7.37 +/- 1.65 x 10-12 cm3/molecule-sec (r2 = 0.99); @ 24C

2.71 +/- 0.20 x 10-12 cm3/molecule-sec (r2 = 0.97); @ 34C

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (30)

Type : air

Light source

Light spectrum : nm

Relative intensity : based on intensity of sunlight

INDIRECT PHOTOLYSIS

Sensitizer : OH

Conc. of sensitizer : 1000000 molecule/cm³

Rate constant : = .0000000000096466 cm³/(molecule*sec)

Degradation : = 50 % after 20 hour(s)

Deg. product

Method : OECD Guide-line draft "Photochemical Oxidative Degradation in the

Atmosphere"

Year : 1990

GLP :

Remark: Calculated value using draft OECD method of Atkinson (1990).

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (81)

Type : air

Light source : other: sun lamp

Light spectrum : nm

Relative intensity: based on intensity of sunlight

Deg. product

Method : other (measured)

Year :

GLP : no data

Test substance

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Under simulated atmospheric conditions and in the presence of 5 ppm NO, the half life of 2-butanol (10 ppm) is 4 hours.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (26)

Remark: 2-Butanol does not contain chromophores that adsorb light at wavelengths

>290 nm. Therefore, direct photolysis will not occur.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

18.01.2002 (20)

3.1.2 STABILITY IN WATER

Type : abiotic t1/2 pH4 : at °C

3. Environmental Fate and Pathways

Id 78-92-2

Date

t1/2 pH7 : at °C **t1/2 pH9** : at °C

Deg. product : Method : Year :

GLP : no data

Test substance :

Result: Hydrolysis will not contribute to the transformation of sBA in aquatic

environments. Hydrolysis of an organic chemical is the transformation process in which a water molecule or hydroxide ion reacts to form a new carbon-oxygen bond, thereby changing the parent chemical. Chemicals that are susceptible to hydrolysis contain functional groups that can be displaced by a nucleophilic substitution reaction. Simple alcohols such as sBA are resistant to hydrolysis because they lack a functional group that is

hydrolytically reactive.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (49)

3.1.3 STABILITY IN SOIL

Remark : If released on land, 2-butanol has the potential to leach

into soil. 2-Butanol also has the potential to volatilize from dry soil.

The estimated log Koc for 2-butanol is 0.84. Therefore, 2-butanol should not absorb significantly to soil or

sediment.

The Koc value was calculated using the equation in Lyman (1982): log Koc = -0.55log S + 3.64 (S, water solubility, in mg/L); water solubility value used

= 125,000 mg/L from Hahn et al. (1986)

[Lyman, W.J. (1982). Adsorption Coefficient for Soils and Sediments. In: Handbook of Chemical Property Estimation Methods. Environmental Behavior of Organic Compounds. 4,1-33. Lyman W.J., Reehl, W.F., and

Rosenblatt, D.H. (eds). McGraw-Hill, New York, NY, USA.]

[Hahn, H-D., Dämbkes, G., and Rupprich, N. (1986). Butanols. In: Gerhartz W (ed), Ullmann's encyclopedia of industrial chemistry, 5th ed, vol A4,

benzyl alcohol to calcium sulfate. VCH, Weinheim, 463-474.]

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
(62) (87)

3.2.1 MONITORING DATA

18.01.2002

Type of measurement : background concentration

Media : surface water

Concentration : Method :

Remark : 2-Butanol has been detected in the Niagara River (Lake

Ontario basin), but not in the western basin of Lake

Ontario.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (43)

ld 78-92-2

Date

Type of measurement

: background concentration

Media

: ai

Concentration

.

Remark

Method

2-Butanol was detected but not quantified in forest air in

the southern Black Forest of Germany.

Source

EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002

(56)

Type of measurement

background concentration

Media

air

Concentration Method

. .

Remark

Low molecular weight alcohols, including 2-butanol, were not detected in air or precipitation samples taken at Tucson, Arizona, and two rural sites 40

km awav.

Source

EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002

(84)

Type of measurement

Media

concentration at contaminated site

Concentration

Method

other: wastewater

Remark

In a comprehensive survey of wastewater from 4000 industrial and publically owned treatment works sponsored by the Effluent Guidelines Division of the U.S. EPA, 2-butanol was identified in discharges of the following industrial category (positive occurences, median concentration in ppb): leather tanning (1; 46.7), petroleum refining (1; 149.3), paint and ink (1; 324.7), organics and plastics (2; 35.4), pesticides manufacture (1; 36.2),

pharmaceuticals (1; 4.6), foundries (1; 41.0), electronics (1; 19.9), mechanical products (3; 90.6), publically owned treatment works (3; 12.4). The highest effluent concentration was 920.1 ppb in the mechanical

products industry.

Source

EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002

(80)

Type of measurement

concentration at contaminated site

Media

other: landfill leachate

Concentration

•

Method

Remark :

2-Butanol was found in landfill leachate from 1 of 5 sites in Connecticut. The concentration in this leachate ranged

from 6.2 to 14.9 ppm.

Source

EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002

(78)

Type of measurement

background concentration

Media

drinking water

Concentration

:

Method

•

Remark

2-Butanol was identified in drinking water from Ottumwa, IA.

Source

EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Id 78-92-2

Date

17.01.2002 (61)

3.2.2 FIELD STUDIES

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type : volatility
Media : water - air

Air : % (Fugacity Model Level I)

Water : % (Fugacity Model Level I)

Soil : % (Fugacity Model Level I)

Biota : % (Fugacity Model Level II/III)

Soil : % (Fugacity Model Level II/III)

Method : other

Year

Result : On the basis of Henry's Law constant, the volatilization half-life of 2-butanol

is estimated to be 120.2 hours in a model river with the follwoing parameters: 1 m deep, 1 m/sec flow rate, and 3 m/sec wind speed.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (86)

3.3.2 DISTRIBUTION

Media : air - biota - sediment(s) - soil - water
Method : Calculation according Mackay, Level I

Year

Remark: Physicochemical data used in the calculation:

Parameter Value w/ Units

Molecular Weight 74.12 Temperature 20 C Log Kow 0.61

Water Solubility 125,000 g/m3 Vapor Pressure 1600 Pa Melting Point -114 C

Result : Using the Mackay Level I calculation, the following

distribution is predicted for 2-butanol:

%Distribution Compartment

 16.28
 Air

 83.68
 Water

 0.03
 Soil

 0.01
 Sediment

0.00 Suspended Sediment

0.00 Biota

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

17.01.2002 (64)

Date

3.4 MODE OF DEGRADATION IN ACTUAL USE

3.5 BIODEGRADATION

Type : aerobic

inoculum : other: Wastewater treatment plant

Contact time : 5 day(s)

Degradation : = 83 (±) % after 5 day(s)

Result

Deg. product

Method : other: APHA (American Public Health Association) Standard Methods, No.

219

Year

GLP : no data

Test substance :

Method: The APHA No. 219 test method, which measures dissolved oxygen, is

similar to the OECD 301D, Closed Bottle biodegradation test procedure.

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result: The ThOD (theoretical oxygen demand) = 2.59 g/g. The measured BOD

(biological oxygen demand) = 2.15 g/g.

The only deviation to the test method was that 0.5 mg/L of allylthiourea was

added to the test medium to prevent nitrification.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition : 10 ml of filtered wastewater was used as an inoculum with 500 ml of test

medium. The source of the inoculum from within the wastewater treatment plant was not stated. The inoculum was not acclimated. There was no

information on controls and replicate test samples.

Test temperature was 20 +/- 1 deg C.

Reliability : (2) valid with restrictions

Although it was cited that a standard method was followed, there is little information in the article confirming that the test was conducted according to the method description. This lack of information supports a reliability

rating of 2.

18.01.2002 (6)

Туре

Inoculum : other: Wastewater treatment plant

Contact time : 50 day(s)

Degradation : (±) % after

Result : Deg. product :

Method : other: Winkler Method

Year :

GLP : no data

Test substance :

Method: The Winkler method, which measures dissolved oxygen, is equivalent to

the OECD 301D, Closed Bottle biodegradation test procedure.

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result : Day % Biodegradation (ThOD)

5 0.0 10 44.2 15 69.2

Id 78-92-2

Date

20 72.3 30 73.2 40 75.4 50 77.0

Source ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition An azide modification of the Winkler test method was used. The source of

the inoculum from within the wastewater treatment plant was not stated. There was no information on controls and replicate test samples. Test

material loading was 2.5 ppm. Test temperature was 20 deg C.

Reliability : (2) valid with restrictions

> There is little information in the article describing how the study was conducted. This lack of information supports a reliability rating of 2. The results were comparable to reported data (Bridie, A.L., C.J.M. Wolff, and M. Winter. 1979. BOD and COD of Some Petrochemicals. Water Research.

13:627-630), which supports the acceptability of this study.

18.01.2002 (59)

Type aerobic

Inoculum activated sludge

Contact time

Degradation $= 98.5 (\pm) \%$ after 5 day(s)

Result

Deg. product

Method other: screening test

Year

GLP no data

Test substance

Remark Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

EXXON CHEMICAL, Limited Fareham, Hampshire Source

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (71)

Type aerobic

Inoculum activated sludge

Contact time

Degradation $= 81.7 (\pm) \%$ after 5 day(s)

Result

Deg. product

Method other: screening test

Year

GLP no data

Test substance

Remark Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

: EXXON CHEMICAL, Limited Fareham, Hampshire Source

other: screening test

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA) (90)

Type aerobic

Inoculum activated sludge

Contact time

Degradation $= 33 (\pm) \%$ after 5 day(s)

Result

Method

17.01.2002

Deg. product

Year

Id 78-92-2

Date

GLP : no data

Test substance

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (28)

Type : aerobic

Inoculum : activated sludge

Contact time

Degradation : = $9.3 \pm 0.3 \pm 0.3$

Result

Deg. product

Method : other: screening test

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (42)

Type : aerobic

Inoculum : other: semi-automatic activated sludge system

Contact time

Degradation : (\pm) % after

Result : other: 98% BOD reduction after 5 days

Deg. product

Method :

Year :

GLP : no data

Test substance: as prescribed by 1.1 - 1.4

Remark: 98% BOD reduction after 5 days

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

14.01.2002 (77)

Type : anaerobic

Inoculum : other: acetate enriched cultures

Deg. product : Method :

Year :

GLP : no data

Test substance : as prescribed by 1.1 - 1.4

Remark: In a long-term study using anaerobic upflow filters, 93%

utilization rate was seen after a 52 day operation.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

14.01.2002 (21)

Type : anaerobic

Inoculum : other: acetate-acclimated methane cultures

Contact time :

Degradation : $100 (\pm) \%$ after 14 day(s)

Id 78-92-2

Date

Result :
Deg. product :
Method :
Year :

GLP : no data

Test substance : as prescribed by 1.1 - 1.4

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

14.01.2002 (92)

3.6 BOD5, COD OR BOD5/COD RATIO

BOD5

Method : other: not specified

Year :

Concentration : related to BOD5 : = 1.87 mg/l concentration : no data

COD

Method : other: not specified

Year

COD : = 2.47 mg/g substance

GLP : no data

RATIO BOD5 / COD

BOD5/COD : = .76

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (57)

3.7 BIOACCUMULATION

BCF : ca. 1.71

Elimination :

Method : other: calculated value

Year

GLP : no data

Test substance :

Remark: A bioconcentration factor of 1.7 is calculated using a 2-butanol log

octanol/water partition coefficient (Kow) of 0.61 as reported by Hansch et al., 1995 (Hansch, C., Leo, A., Hoekman, D. 1995. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American Chemical

Society, Washington, DC, USA.).

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA) 18.01.2002 (16)

3.8 ADDITIONAL REMARKS

Memo : The volatilization half-life in a model river is estimated to be 3.5 days at 25

degrees C.

Id 78-92-2

Date

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (86)

Date

4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type : flow through

Species: Pimephales promelas (Fish, fresh water)

Exposure period : 96 hour(s)
Unit : mg/l

LC50 : = 3670 measured/nominal

Limit test

Analytical monitoring : yes

Method : other: no data

Year : 1985 GLP : no data

Test substance : as prescribed by 1.1 - 1.4

Method : The test method used in this study is similar to the OECD 203 test

guideline.

6,667

Trimmed Spearman-Karber Method (Hamilton, M.A., R.C. Russo, and R.V. Thurston. 1977. Trimmed Spearman-Karber Method for Estimating Median Lethal Concentrations in Toxicity Bioassays. Environ. Sci. Technol. 11:714-

719. Correction, 12:417, 1978.)

Remark : Purity: 99%

Result : 96-hour LC50 = 3,670 mg/L (95% CI 3,380 to 3,990) based upon average,

corrected measured values

Analytical method used was Gas-Liquid Chromatography
Measured Fish Total

Conc. (mg/L) Mortality (@96 hrs)*

Control 0

1,018 0

1,839 0

2,630 0

4,258 16

* 20 fish added at test initiation

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition: Treatment solutions were prepared by diluting a 3.7 g/L stock solution.

20

Nominal sBA treatment levels were 1.05, 1.61, 2.48, 3.82, 5.88mg/L, which

measured 1.02, 1.84, 2.63, 4.26 and 6.67mg/L, respectively.

Control/dilution water was EPA Duluth laboratory water. Twenty fish were tested per treatment and control. Tank volume = 2L. Control and treatment

solution flow rate was equivalent to 18 chamber volumes per day. Mean test parameters were as follows: temperature = 24.4 deg. C; dissolved oxygen = 7.5 mg/L; pH = 7.8; fish age = 30 days; fish mean wt. = 0.09 g; fish mean length = 18.9 mm; fish loading = 0.0.90 g/L. Organism supplier was U.S. EPA Environmental Research Lab, Duluth, MN, USA.

Reliability : (2) valid with restrictions

18.01.2002 (41)

Type : static

Species: Carassius auratus (Fish, fresh water)

Exposure period : 24 hour(s)
Unit : mg/l

LC50 : = 4300 measured/nominal

Limit test

Analytical monitoring : yes

Method : other: APHA (American Public Health Association) Standard Methods

Year

GLP : no data

Test substance :

Date

Method : Interpolation from a graph of log concentration values versus percent

mortality (APHA, 1971).

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition : A series of treatment solutions were prepared, but their concentrations

were not reported. Ten fish were tested per treatment. Tank volume = 25L.

Treatment solutions were aerated during the test. Treatment

concentrations were analytically verified at test start and termination, but the method of analysis was not reported. Reported results were not distinguished as being based on nominal or measured values and there is

no control information.

Test parameters were as follows: temperature = 20+/-1 deg. C; dissolved oxygen did not fall below 4 mg/L; initial pH = 7.0; fish average wt. = 3.3+/-

1.0 g; fish average length = 6.2+/-0.7 cm.

Reliability : (2) valid with restrictions

It is unclear from the article as to whether the reported result is based on nominal or measured values. This lack of information, in conjunction with the absence of selected test parameters and results (i.e., exposure concentrations, mortality), supports a reliability rating of 2. The reported value in this study is comparable to a reported fathead minnow acute value for sBA: 24-hour LC50 = ~4,000 mg/L (Geiger D.L. et al. 1986. Acute Toxicities of Organic Chemicals to Fathead Minnows (Pimephales promelas), Vol. 3. Center for Lake Superior Environmental Studies.

University of Wisconsin-Superior, WS, USA.).

17.01.2002 (7)

Type : static

Species : Leuciscus idus (Fish, fresh water)

Exposure period : 48 hour(s)
Unit : mg/l

LC50 : = 3520 measured/nominal

Limit test

Analytical monitoring : no

Method : other: not specified

Year :

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (55)

Type : static

Species: Petromyzon marinus

Exposure period : 24 hour(s)

Unit

Limit test :

Analytical monitoring : no

Method : other: not specified

Year :

GLP : no data

Test substance :

Method : Water used in the study came from Hammond Bay of Lake Huron and

taken from a point source that was located 250 feet offshore at a depth of approximately nine feet. Test systems were 10L glass jars that contained 5L of lake water. Test systems were aerated. Up to six organisms were

Date

added to each test system. Larval lampreys (Petromyzon marinus) were collected by means of an electric shocker in the Ocqueoc River, Presque Isle County, Michigan, USA, and were held in running water in aquaria and samll "races" under conditions which simulated their natrual stream habitat.

Water pH ranged from 7.5 to 8.2; water temperature was 55+/-1 degrees F; dissolved oxygen ranged from 8.6 to 13.7 ppm; and free CO2 ranged from 5.0 to 9.0 ppm. Treatment solutions used were 5.0, 1.0, and 0.1 ppm. A control system that only contained lake water was included. It was not stated whether the units were presented as a volume or weight.

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result : The highest treatment level tested was 5 ppm. No effects were observed at

this level to laral lamprey.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

18.01.2002 (4)

Type

Species : other: fish
Exposure period : 96 hour(s)
Unit : mg/l

LC50 : = 1113 calculated

Method : other: ECOSAR Computer Model

Year :

GLP :

Test substance :

Remark : Test Type: Acute Fish Toxicity Calculation Source : ExxonMobil Biomedical Sciences, Inc., NJ, USA

Test condition: A log Kow (octanol/water partition coefficient) value and a chemical

structure are needed to complete the calculation with this model. The log Kow value used was 0.61. The SMILES (Simplified Molecular Input Line Entry System) structure used with the model was: OC(CC)C. Additional data calculated by the model include molecular weight, 74.12, and water

solubility, 8,625 mg/L (@25C).

The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American

Chemical Society. Washington, DC, USA.

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

17.01.2002 (18)

4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

Type : static

Species : Daphnia magna (Crustacea)

Exposure period : 48 hour(s)
Unit : mg/l

EC50 : = 4227 measured/nominal

Analytical monitoring : no

Method : other: German Institute of Standardization, DIN 38412, Part II, Daphnia

Short-Time Test

Year :

GLP : no data

Test substance : as prescribed by 1.1 - 1.4

Method : None applied. The EC50 was calculated arithmetically from the

concentration/effect ratio.

Date

The test method used in this study is similar to the OECD 202 test

guideline.

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result : 48-hour EC50 = 4,227 mg/L (95% CI 1,859 to 7,143) based on nominal

values

The authors considered the test valid because fewer than 10% of the organisms in the control were immobile, the pH value was not below 7.0,

and the DO was not below 4.0 mg/L at test termination.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition: The test procedure used daphnids 6 to 24 hours old. Organisms were not

fed during the test. The treatment medium was defined and had a total hardness of 2.4 mmol/L, a calcium to magnesium ratio of 4:1, a sodium to potassium ratio of 10:1, and an initial pH value of 8.0+/-0.2. The test

solution temperature was 20°C.

The test systems used 50 ml beakers. Treatment solutions and controls were prepared in duplicate. The organism loading rate was no less than one animal per 2 ml test medium. 20 organisms were tested per treatment and control. Treatment solutions were prepared from a stock sBA solution. Dissolved oxygen (DO) and pH were measured at test termination although

specific values were not supplied for this test.

Reliability : (2) valid with restrictions

Although the method was described in the article, data were not provided on the test parameters or results from individual treatment and control solutions. This lack of information supports a reliability rating of 2.

18.01.2002 (58)

Type : static

Species : Daphnia magna (Crustacea)

Exposure period : 24 hour(s)
Unit : mg/l

LC50 : = 3750 measured/nominal

Analytical monitoring : no

Method : other: not specified

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (9)

Type : static

Species: other: Tetrahymena pyrifomis (Ciliate)

Exposure period : 48 hour(s)
Unit : mg/l

EC50 : = 3196 measured/nominal

Analytical monitoring : no data

Method : other: Population Growth Impairment Test

Year :

GLP : no data

Test substance :

Method : Probit analysis procedure using Statistical Analysis System (SAS) software

(SAS Institute, 1989).

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result : 48-hour EC50 for growth = 3,196 mg/L based on nominal values

Population density (growth) was measured spectrophotometrically as

Date

absorbance at 540 nm.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition: Axenic cultures of Tetrahymena pyrifomis were used in the test. Only 48-

hour population densities were measured. Treatment solutions and controls were prepared in duplicate. Treatment solutions were prepared from a

stock sBA solution.

Reliability : (2) valid with restrictions

A non-standardized method was referenced in the article. Data were not provided on the test parameters or results from individual treatment and control solutions. This lack of information supports a reliability rating of 2.

17.01.2002 (79)

Туре

Species : other: Daphnid : 48 hour(s) Unit : mg/l

LC50 : = 1084 calculated

Method : other: ECOSAR Computer Model

Year :

GLP : Test substance :

Remark: Test Type: Acute Daphnid Toxicity Calculation

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition: A log Kow (octanol/water partition coefficient) value and a chemical

structure are needed to complete the calculation with this model. The log Kow value used was 0.61. The SMILES (Simplified Molecular Input Line Entry System) structure used with the model was: OC(CC)C. Additional data calculated by the model include molecular weight, 74.12, and water

solubility, 8,625 mg/L (@25C).

The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American

Chemical Society. Washington, DC, USA.

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

17.01.2002 (18)

4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE

Species : Scenedesmus quadricauda (Algae)

Endpoint

Exposure period : 7 day(s)
Unit : mg/l

TT : = 95 measured/nominal

Limit test

Analytical monitoring : no

Method : other: 7-Day Cell Multiplication Inhibition Test

Year :

GLP : no data

Test substance :

Method: None applied. The toxicity threshold (TT) was determined graphically by

plotting the highest non-toxic concentration versus its mean extinction value against the lowest toxic concentration versus its mean extinction value and calculating the toxicant concentration at 3% below the no effect

level.

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%. Test Type: Static Toxicity Test

Result: 7-day TT (toxicity threshold) for growth = 95 mg/L based on nominal

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values.

The TT value for growth is calculated by identifying the treatment level that is greater or equal to 3% below the treatment level that did not exhibit toxic effects as measured by the extinction of primary light of monochromatic

radiation at 578 nm.

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA Source

Treatment solutions were prepared by diluting a stock sBA solution. **Test condition**

Testing was conducted in metal capped, 300 ml Erlenmeyer flasks containing 50 ml of treatment solution. Treatment solutions contained sBA, cells, double distilled water, and a sterile, defined nutrient medium. The control solution contained nutrient medium, to which sterile double distilled water was added. Growth inhibition measurements were only determined

on day 7.

Cell growth was determined by using a turbidimetric procedure that measured primary light extinction (monochromatic radiation at 578 nm)

through a cell suspension of 10 mm thickness.

(2) valid with restrictions Reliability

> Although a non-standardized method was described in the article, data were not provided on the test parameters, replication, or results from individual treatment and control solutions. This lack of information supports

a reliability rating of 2.

17.01.2002 (13)

Species other algae: green alga

Endpoint

Exposure period 96 hour(s) Unit mg/l

EC50 = 625 calculated

Method other: ECOSAR Computer Model

Year **GLP**

Test substance

Remark Test Type: Green Alga Toxicity Calculation

96-hour EC50 = 625 mg/LResult

ExxonMobil Biomedical Sciences, Inc., NJ, USA Source

Test condition A log Kow (octanol/water partition coefficient) value and a chemical

> structure are needed to complete the calculation with this model. The log Kow value used was 0.61. The SMILES (Simplified Molecular Input Line Entry System) structure used with the model was: OC(CC)C. Additional data calculated by the model include molecular weight, 74.12, and water

solubility, 8,625 mg/L (@25C).

The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American

Chemical Society. Washington, DC, USA.

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

17.01.2002 (18)

Species

Endpoint other: green alga **Exposure period** 96 hour(s)

Unit mg/l

ChV* = 28 calculated

Method other: ECOSAR Computer Model

Year

GLP Test substance

Result *Chronic value

Source ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition A log Kow (octanol/water partition coefficient) value and a chemical

Date

structure are needed to complete the calculation with this model. The log Kow value used was 0.61. The SMILES (Simplified Molecular Input Line Entry System) structure used with the model was: OC(CC)C. Additional data calculated by the model include molecular weight, 74.12, and water

solubility, 8,625 mg/L (@25C).

The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American

Chemical Society. Washington, DC, USA.

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

18.01.2002 (18)

Species: Microcystis aeruginosa (Algae, blue, cyanobacteria)

Endpoint

Exposure period : 192 hour(s)
Unit : mg/l

NOEC : = 312 measured/nominal

Limit test

Analytical monitoring : no data

Method : other: not specified

Year :

GLP : no data

Test substance

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

18.01.2002 (8)

Species : Chlorella pyrenoidosa (Algae)

Endpoint

Exposure period

Unit : mg/l

NOEC : = 8900 measured/nominal

Limit test

Analytical monitoring : no data

Method : other: not specified

Year

GLP : no data

Test substance

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

18.01.2002 (54)

4.4 TOXICITY TO MICROORGANISMS E.G. BACTERIA

Type : aquatic

Species : Pseudomonas putida (Bacteria)

Exposure period : 16 hour(s)
Unit : mg/l

ECO : ca. 500 measured/nominal

NOAEL : = 500 Analytical monitoring : no data

Method : other: total biomass

Year

GLP : no data

Test substance

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Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

18.01.2002 (12)

Type

Species : Bacillus subtilis (Bacteria)

Exposure period

Unit : mg/l

EC50 : = 1630 measured/nominal

Analytical monitoring : no data

Method :

Year

GLP : no data

Test substance :

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

18.01.2002 (36)

Type : aquatic

Species : Chilomonas paramaecium (Protozoa)

Exposure period : 48 hour(s)
Unit : mg/l

NOEL : = 745 measured/nominal

Analytical monitoring : no data

Method : other: total biomass

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

18.01.2002

Type : aquatic

Species : Entosiphon sulcatum (Protozoa)

Exposure period : 72 hour(s)
Unit : mg/l

NOEL : = 1282 measured/nominal

Analytical monitoring : no data

Method : other: total biomass

Year :

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

18.01.2002 (11)

Type : aquatic

Species : Uronema parduzci (Protozoa)

Exposure period : 20 hour(s)
Unit : mg/l
NOEC : = 1416
Analytical monitoring : no data

Method : other: total biomass

Date

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

17.01.2002 (10)

4.5.1 CHRONIC TOXICITY TO FISH

Species: other: fish

Endpoint :

Exposure period : 30 day(s)
Unit : mg/l

ChV* : = 115 calculated

Method : other: ECOSAR Computer Model

Year : GLP : Test substance :

Remark: Test Type: Chronic Fish Toxicity Calculation

* Chronic Value

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition : A log Kow (octanol/water partition coefficient) value and a chemical

structure are needed to complete the calculation with this model. The log Kow value used was 0.61. The SMILES (Simplified Molecular Input Line Entry System) structure used with the model was: OC(CC)C. Additional data calculated by the model include molecular weight, 74.12, and water

solubility, 8,625 mg/L (@25C).

The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American

Chemical Society. Washington, DC, USA.

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

17.01.2002 (18)

4.5.2 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES

Species : other: Daphnid

Endpoint

Exposure period : 16 day(s)
Unit : mg/l

EC50 : = 30 calculated

Method : other: ECOSAR Computer Model

Year : GLP : Test substance :

Remark : Test Type: Chronic Daphnid Toxicity Calculation

Source: ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition : A log Kow (octanol/water partition coefficient) value and a chemical

structure are needed to complete the calculation with this model. The log Kow value used was 0.61. The SMILES (Simplified Molecular Input Line Entry System) structure used with the model was: OC(CC)C. Additional data calculated by the model include molecular weight, 74.12, and water

solubility, 8,625 mg/L (@25C).

The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.

Date

Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American

Chemical Society. Washington, DC, USA.

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

17.01.2002 (18)

4.6.1 TOXICITY TO SEDIMENT DWELLING ORGANISMS

4.6.2 TOXICITY TO TERRESTRIAL PLANTS

Species: other terrestrial plant: Solanum tuberosum L.

Endpoint :

Exposure period Unit Method

Year

GLP : no data

Test substance :

Method : stage at application: seedling, on intact plant, in

environmental chamber, application by fumigation: 2 cm3/l.

Remark : Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result : 20% respiration increase, on tuber

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (66)

Species: other terrestrial plant: wheat

Endpoint

Exposure period :

Unit
Method

Year

GLP : no data

Test substance

Method : Stage at application: seedling, on excised organ; addition

to growth medium, dose: 4 x 10E-2 M

Remark : Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result: 50% decrease in number of cells

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA) 17.01.2002 (45)

Species : other terrestrial plant: wheat

Endpoint :

Exposure period : Unit : Method : Year :

GLP : no data

Test substance

Method : Stage at application: seedling, on excised organ; addition

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to growth medium, dose: 5 x 10 E-2 M.

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result: 50% size decrease of cells

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (44)

Species: other terrestrial plant: Lupinus albus

Endpoint

Exposure period : 1 day(s)

Unit Method Year

GLP : no data

Test substance :

Method: Stage at application: seedling, on intact plant; addition to

growth meduim, in culture flask, dose = 1%.

Remark : Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Result: 73% size decrease of root

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (63)

Species : Lactuca sativa (Dicotyledon)
Endpoint : other: seed germination

Exposure period

Unit : mg/l **EC50** : = 650

Method : other: not indicated

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (91)

Species: other terrestrial plant: Cucumis sativus

Endpoint : other: seed germination

Exposure period

Unit : mg/l **NOEC** : < 50375

Method : other: not specified

Year

GLP : no data

Test substance :

Remark: Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

17.01.2002 (91)

Date

4.6.3 TOXICITY TO SOIL DWELLING ORGANISMS

Type

Species: other: earthworm

Endpoint : mortality
Exposure period : 14 day(s)
Unit : other: ppm

LC50 : = 1222 calculated

Method : other: ECOSAR Computer Model

Year GLP Test substance

Remark: Test Type: Earthworm Toxicity Calculation

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition: A log Kow (octanol/water partition coefficient) value and a chemical

structure are needed to complete the calculation with this model. The log Kow value used was 0.61. The SMILES (Simplified Molecular Input Line Entry System) structure used with the model was: OC(CC)C. Additional data calculated by the model include molecular weight , 74.12, and water

solubility, 8,625 mg/L (@25C).

The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995. Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American

Chemical Society. Washington, DC, USA.

Reliability : (2) valid with restrictions

Rated 2 for reliability due to calculated value.

17.01.2002 (18)

4.6.4 TOX. TO OTHER NON MAMM. TERR. SPECIES

Species: other: Xenopus laevis (Clawed Frog)

Endpoint : mortality
Exposure period : 48 hour(s)
Unit : other: mg/L

LC50 : = 1530 measured/nominal

Method : other: not specified

Year

GLP : no data

Test substance :

Method: Median lethal concentration was calculated as a projection from the least

square linear regression on log transformed nominal concentration data

and probit transformed percent effect data.

Remark: Test Type: Static Acute Toxicity Test

Analytical Monitoring: No

Results based on nominal treatment values.

Information on sBA purity is not available; assumed to have used a

commercial grade of >/= 99%.

Source : ExxonMobil Biomedical Sciences, Inc., New Jersey, USA

Test condition: The test procedure used 3 to 4 week old larvae. The organisms were

exposed in covered all-glass aquaria containing 1L of Dutch Standard Water at a temperature of 20+/-1C. The test material was added once at the beginning of the study. Five concentration levels were evaluated with 10 organisms per level. The 5 levels had a factorial difference of 1.5.

Mortality was only recorded after 48 hours.

The test species, Xenopus laevis, was identified in the article as a clawed toad. However, it is correctly identified as a clawed frog in this robust

summary.

Reliability : (2) valid with restrictions

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The test procedure was generally described in the article. Data were not provided on the test parameters or results from individual treatment and control solutions. This lack of information supports a reliability rating of 2.

18.01.2002

(24)

- 4.7 BIOLOGICAL EFFECTS MONITORING
- 4.8 BIOTRANSFORMATION AND KINETICS
- 4.9 ADDITIONAL REMARKS

Date

5.0 TOXICOKINETICS, METABOLISM AND DISTRIBUTION

In Vitro/in vivo :
Type :
Species : rat

Number of animals

Males : Females :

Doses

Males : (See Remark)

Females

Vehicle : other: Aqueous Solution

Route of administration : other: Oral Gavage / i.v.

Exposure time : Product type guidance : Decision on results on acute tox. tests : Adverse effects on prolonged exposure :

Half-lives : 1st: 2nd: 3rd.

Toxic behaviour : Deg. product :

Method : other: Experimental Model Development

Year : 1981 GLP : no data

Test substance : other TS: 2-Butanol (sec-Butanol; sBA) and metabolites 2-Butanone

(Methyl ethyl ketone; MEK), 3-Hydroxy-2-Butanone (3H-2B), and 2,3-Butanediol (2,3-BD) (CAS Nos. 78-92-2 (sBA); 78-93-3 (MEK))

Method : Statistical Methods

Student's t-test was used for statistical evaluation of differences between two means. In the pharmacokinetic model, the differential equations were solved numerically by a Hammings Predictor-Correction method with a

Runge-Kutta starter.

Remark: Dietz et al. (1981) selected doses for MEK and sBA based on the earlier

pharmacokinetic analysis (Traiger and Bruckner, 1976) that established approximately 96% of an administered dose of sBA was oxidized in vivo to MEK. Thus 1960 mg/kg (2.1 ml/kg) of MEK estimates the amount of MEK formed in vivo from a dose of 1776 mg/kg (2.2 ml/kg) of sBA. Male rats were given oral doses of sBA, MEK, 2-3 B-D (or i.v.), or 3H-2B (i.v.) and serial blood samples were collected for up to 30 hours following treatment.

Test Type: Metabolism and Pharmacokinetic Evaluation

Strain: Sprague-Dawley

Dose Groups/Concentrations: sBA: 2.2 ml/kg or 1776 mg/kg as a 22% aqueous solution (oral). MEK: 2.1ml/kg or 1690 mg/kg as a 21% aqueous solution (oral). 2,3-BD: 0.68 ml/kg or 676 mg/kg as a 6.8 % aqueous solution (oral or i.v.). 3H-2B: 400 or 800 mg/kg as a 60% aqueous solution

(i.v.)

Frequency of Treatment: Single Dose (oral or i.v.)

Duration of Test: 30 Hours

Sex: Male

Number/Dose Group: Not Identified

Result: In the sBA-treatment phase of the study, blood concentrations of sBA and

its metabolites MEK, 3H-2B, and 2,3-BD were measured. Blood sBA concentrations of 0.59 mg/ml peaked at 2 hours and declined to less than 0.05 mg/ml at 16 hours. As the sBA concentration fell, the metabolite concentrations of MEK, 3H-2B, and 2,3-BD concentrations rose to

maximums at 8, 12, and 18hr, respectively. The peak concentration of MEK was 0.78 mg/ml, while that of 2,3-BD was 0.21 mg/ml. 3H-2B reached a

Id 78-92-2 5. Toxicity

Date

peak concentration of 0.04 mg/ml. Total AUC values for sBA, MEK, 3H-2B, and 2,3-BD were 3254 ± 258 , 9868 ± 566 , 443 ± 93 , and 3167 ± 503 mg-

hr/l, respectively.

Following an oral dose of MEK, blood concentrations of MEK and its metabolites sBA, 3H-2B, and 2,3-BD were measured. Blood MEK concentrations of 0.95 mg/ml peaked at 4 hours and declined to less than 0.07 mg/ml at 18 hours. As the MEK concentration fell, the end metabolite 2.3-BD rose to a maximum concentration of 0.26 mg/ml at 18 hours. Peak concentrations of sBA and 3H-2B were 0.033 and 0.027 mg/ml, respectively. These were detected at 6 and 8 hr after the MEK administration. Total AUC values for MEK, sBA, 3H-2B, and 2,3-BD were $10,899 \pm 824,414 \pm 38,382 \pm 38,$ and 3863 ± 238 mg-hr/l, respectively.

2-Butanol (sec-Butanol; sBA) and metabolites 2-Butanone (Methyl ethyl

ketone; MEK), 3-Hydroxy-2-Butanone (3H-2B), and 2,3-Butanediol (2,3-

BD) (CAS Nos. 78-92-2 (sBA); 78-93-3 (MEK))

Purity not identified.

In the development of the model, the authors confirmed that a limited Conclusion

> amount of sBA could be recovered as glucuronide in the urine. Most of the alcohol appears to undergo oxidation via alcohol dehydrogenase to its corresponding ketone, MEK. The results showed a limited amount of the ketone undergoes a backward reduction to its parent alcohol, sBA. 3H-2B and 2,3-BD are found to be common metabolites of sBA and MEK. The model was able to simulate blood concentrations and elimination of all 4 compounds after oral administration of sBA, and the results after i.v. of 3H-2B and 2,3-BD. AUC analysis suggested that the quantities of 3H-2B and 2,3-BD formed from oral doses of sBA and MEK are comparable. The results supported the estimation in that no significant difference in the AUC of MEK was observed after dosing with either 1776 mg/kg of sBA or 1690 mg/kg of MEK (10,899 \pm 824 vs. 9868 \pm 566 mg-hr /liter, respectively).

Reliability (1) valid without restriction

> No circumstances occurred that would have affected the quality or integrity of the data. Test procedures were in accordance with generally accepted

scientific standards and described in sufficient detail.

01.06.2006 (25)

In Vitro/in vivo Type

Species guinea pig

Number of animals

Test substance

Males Females

Doses

Males 450 mg/kg body weight

Females

Vehicle other: Corn oil (25% solution)

Route of administration : i.p. **Exposure time**

Product type guidance Decision on results on acute tox, tests Adverse effects on prolonged exposure

1st: Half-lives $2^{n\dot{d}}$. $3^{\text{rd.}}$

Toxic behaviour Deg. product

Method other: Experimental

Year 1976 **GLP** no data

Test substance other TS: 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)

Male guinea pigs ranging in weight from 250 - 450 grams were given a Remark

single i.p. dose of 450mg/kg MEK (study also investigated MnBK and MiBK

Date

- data not addressed here). Blood was collected by heart puncture from 4 animals at each of the following times after dose administration: 1, 2, 4, 6, 8, 12, and 16 Hr. Only 1 sample was collected from each guinea pig. Serum was separated and refrigerated until assayed within 48 Hr. The concentrations of the ketones and their metabolites were measured in duplicate by direct on-column injection of undiluted serum into a Varian 2100 Gas Chromatograph equipped with a flame ionization detector. Ketones and metabolites were quantitated from calibration curves prepared from pure standards. Greater than 90% of each ketone was distributed in the plasma fraction. Half-lives were estimated by extrapolating the linear portion of the decay curve to zero time.

Test Type: Metabolism and Pharmacokinetic Evaluation

Frequency of Treatment: Single injection

Duration of Test: 16 Hours

Number/Dose Group: 4 at each blood - sampling interval

Result : MEK: 2-Butanol, 3-hydroxy-2-butanone, and 2,3-butanediol were identified

as metabolites in the serum of guinea pigs injected i.p. with MEK. The half-life of MEK in serum was 270 min. The clearance time of MEK was 12 Hr. 2,3-Butanediol was cleared in 16 hours, as were the other two metabolites. The metabolism was described as oxidation via hydroxylation of the ? -1 carbon forming 3-hydroxy-2-butanone as the metabolite of MEK. Reduction occurred at the carbonyl group as expected forming the secondary alcohol,

2-butanol from MEK.

Test substance : 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)

Purity: 98% with traces of 2-butanol

Conclusion: This study showed the initial metabolism of MEK follows both oxidative and

reductive pathways to produce 3-hydroxy-2-butanone, 2,3-butanediol and

2-butanol.

Reliability : (1) valid without restriction

No circumstances occurred that would have affected the quality or integrity of the data. Test procedures were in accordance with generally accepted

scientific standards and described in sufficient detail.

01.06.2006 (27)

In Vitro/in vivo : Type :

Species : human

Number of animals

Males : 9 Females :

Doses

Males : 200 ppm (8.2mmol/m3)

Females

Vehicle : other: none

Route of administration : inhalation

Exposure time : Product type guidance : Decision on results on acute tox. tests : Adverse effects on prolonged exposure :

Half-lives : 1st: 2nd.

2rd:

Toxic behaviour : Deg. product :

Method : other: Experimental

Year : 1988 GLP : no data

Test substance: other TS: 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)

Remark: Nine healthy male volunteers, 18-34 years of age (mean 23.7), weight 65-

81 kg (mean 74.3), height 172-190 cm (mean 181.8) and calculated

surface area 1.81 - 2.09 m2 (mean 1.97) were exposed for 4 Hr to 200ppm

Date

MEK on 2 separate days at least 1 week apart. Venous blood samples were collected at 1-hr intervals during exposure and over 120 to 210 minutes post-exposure. In a supplementary study, follow-up elimination blood samples were collected in 2 persons until the next morning. One of the exposures constituted sedentary activity and the other encompassed three 100 W ergometric exercise periods over minutes 5-15, 95-105 and 225-235 during a total exposure of 240 minutes. Exhaled air samples were collected with a 2-way respirator mouthpiece into 4-liter polyester laminated aluminum-foil bags and analyzed immediately. Samples were collected at one-hour intervals during exposure and over 120-210 minutes thereafter. Urine samples were obtained at 2-hour intervals during the exposure day and in separate samples until the next morning. Whole venous blood was analyzed by gas chromatography, as were air samples. Peaks were compared to calibration curves prepared of known blood or air concentrations of MEK. 2-3BD in urine was analyzed according to a modification of the method of Robinson and Reive. The 2 peaks in the chromatograph (d,l-forms and meso-form) were summed for calculations. Test Type: Metabolism and Pharmacokinetic Evaluation Frequency of Treatment: 2 Exposure periods at least 1 wk between Duration of Test: 4 Hr. Exposure

Result

Pulmonary retention of MEK in the lungs remained constant throughout exposure with or without exercise. Relative uptake was $53\% \pm 2\%$. Estimated mean total pulmonary uptakes were 11.4 mmol (ventilation volume 11 liters/min at rest) and 14.3 mmol (ventilation volume 35 liters/min during the exercise). Blood MEK concentrations increased rapidly during the first hour of exposure (markedly faster when associated with exercise). Thereafter, concentrations increased slowly and linearly through 4 hr with sedentary activity and steeply during the exercise period at the end of the 4 hr exposure period. Two elimination phases were detected for MEK in blood. The calculated half-time for the faster phase of elimination (0-45 min post-exposure) was 30 min and about 81 min for the slower phase (60-320 min post-exposure). Owing to remarkable solubility of MEK, only 2-3% of absorbed dose was eliminated by exhalation. Urinary excretion of unchanged MEK was 0.1% and excretion of the metabolite 2H3B was about 0.1%. The authors speculated that the greater part of absorbed MEK is probably converted to products of intermediary metabolism, e.g., to acetate or acetoacetate via 3H2B.

Test substance

2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)

Purity: Analytical Grade

(1) valid without restriction

Conclusion

Two elimination phases were detected for MEK in blood. The T½ for the faster phase of elimination was 30 min and about 81 min for the slower phase. 2-3% of absorbed dose of MEK was eliminated by exhalation. Urinary excretion of unchanged MEK was 0.1% and excretion of the metabolite 2H3B was about 0.1%.

Reliability

No circumstances occurred that would have affected the quality or integrity of the data. Test procedures were in accordance with generally accepted

scientific standards and described in sufficient detail.

01.06.2006 (60)

5.1.1 ACUTE ORAL TOXICITY

Type : LD50

Value : = 6500 mg/kg bw

Species : Strain : Sex : Number of animals : Vehicle : Doses :

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Method : other

Year

GLP : no data

Test substance: as prescribed by 1.1 - 1.4

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

21.10.1992 (34)

Type : LD50

Value : = 2054 - 2328 mg/kg bw

Species : rat

Strain : Fischer 344
Sex : male/female

Number of animals : 10

Vehicle : other: None

Doses

Method : other: Experimental (Non-regulatory)

Year : 1986 **GLP** : yes

Test substance: as prescribed by 1.1 - 1.4

Remark: Fischer 344 rats were in the age range of 55-65 days when purchased and

were 9 -11 weeks old when the study commenced. All animals were fasted for 18 hours prior to dosing and doses were altered by varying the volume dispensed from the syringe. Body weights were recorded on Days 1, 7 and 14. All surviving rats gained weight by the end of the 14-day observation

period. The LD50's were calculated using probit analysis

Route of Administration: Oral Gavage

Doses: 950 - 2400 mg/kg

Volume Administered: Single dose volume varied by body weight

Post Dose Observation Period: Daily for 14 Days

Purity: 99.5%

Result : 2054 mg/kg (95% fiducial limits, 1283 - 4018 mg/kg) males,

2328 mg/kg (95% fiducial limits, 1470 - 5428 mg/kg) females, and 2193 mg/kg (95% fiducial limits, 1608 - 4146 mg/kg) combined.

Conclusion: The acute oral LD50 for sBA in Fischer 344 rats is 2193 mg/kg.

Reliability : (1) valid without restriction

1 - Reliable study without restrictions. No circumstances occurred that

would have affected the quality or integrity of the data.

10.01.2002 (72)

Type : LD50

Value : = 5730 - 7320 mg/kg bw

Species : rat

Strain : other: Carworth-Wistar

Sex : male Number of animals : 5

Vehicle : other: Unknown (water, corn oil, or a 1% solution of 3,9-diethyl-6-

tridecanol sulfate (Tergitol Penetrant 7)

Doses

Method : other: Experimental (Non-regulatory)

Year : 1954 **GLP** : no

Test substance : as prescribed by 1.1 - 1.4

Remark : Route of Administration: Oral Gavage

Doses: Logarithmic series

Dose Volume Administered: Single dose; 1-10 ml/rat

Post Dose Observation Period: 14 Days

The animals weighed 90 - 120 grams and were not fasted prior to dosing. The most probable LD50 value and the fiducial range were estimated by

5. Toxicity ld 78-92-2
Date 01.06.2006

the method of Thompson using the tables of Weil.

Result : 6.48 g/kg (5.73 - 7.32)

Conclusion : The acute oral LD50 for sBA in Carworth-Wistar rats is 6.48 g/kg (5.73 -

7.32)

Reliability : (2) valid with restrictions

Smyth, H. F. Jr., Carpenter, C. P., Weil, C. S., and Pozzani, U. C. (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med, 10:61-68.

10.01.2002 (83)

Type : LD50

Value : = 4900 mg/kg bw

Species : rabbit

Strain :
Sex :
Number of animals :
Vehicle :
Doses :

Method : other: not specified

Year : 1972 GLP : no data

Test substance : as prescribed by 1.1 - 1.4

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

21.10.1992

Type : LD50

Value : = 4890 mg/kg bw

Species : rabbit

Strain : other: Unknown
Sex : male/female

Number of animals

Vehicle : no data

Doses

Method : other: Experimental (Non-regulatory)

Year : 1925 **GLP** : no

Test substance: as prescribed by 1.1 - 1.4

Remark : Route of Administration: Oral Gavage

Number of Animals/Dose: Unknown

Doses: 14 or 66 mmol/kg; 1037 or 4890 mg/kg

Dose Volume Administered: < 50 ml Post Dose Observation Period: 24 Hours

Ten to 35 rabbits weighing between 1.5 and 2.5 kg were given "calculated" quantities of sBA at doses described as the "Narcotic Dose " or "Certain

Lethal Dose" by oral gavage and observed for 24 hours.

The cited LD50 [66 mmol/kg, or 4890 mg/kg] is really the Certain Lethal Dose presented in the paper by Munch and Schwartze (1925). The Munch paper (1972) presents the rabbit data reported earlier in 1925 plus tadpole data. Because of the inconsistency in the definition of LD50 and Certain

Lethal Dose, a CoR of 3 is assigned to both references.

Result : 4890 mg/kg

Conclusion : The reported LD50 was 66 mmol/kg (4890 mg/kg).

Reliability : (3) invalid

3 - Not Reliable. Study is pre-GLP and documentation insufficient for

assessment

10.01.2002 (67) (68)

Date

5.1.2 ACUTE INHALATION TOXICITY

Type : LC50

Value : = 16000 ppm

Species : rat

Strain : other: Carworth-Wistar

Sex : male Number of animals : 6

Vehicle : other: None

Doses

Exposure time : 4 hour(s)

Method : other: Experimental (Non-regulatory)

Year : 1954 **GLP** : no

Test substance : as prescribed by 1.1 - 1.4

Remark: Route of Administration: Inhalation

Observation Period: Up to 14 days

The study consisted of exposing six male albino rats to a flowing stream of air approaching saturation with sBA vapors. The stream was prepared by proportioning pumps and nominal concentrations (not confirmed by analytical methods) were recorded. The study end point was the concentration yielding a fractional mortality among six rats within 14 days.

Result : 16,000 ppm sBA was lethal to 5 of 6 rats within 14 days

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Conclusion: Five of six exposed rats died within 14 days.

Reliability : (2) valid with restrictions

2- Reliable with restrictions. Pre-GLP study documented, meets generally

accepted scientific principles, acceptable for assessment.

11.01.2002 (83)

Species : rat

Strain : other: Carworth-Wistar

Sex : male Number of animals : 6

Vehicle : other: None

Doses

Exposure time : 4 hour(s)

Method : other: Experimental (Non-regulatory)

Year : 1951 GLP : no

Test substance: as prescribed by 1.1 - 1.4

Remark : Route of Administration: Inhalation

Observation Period: Up to 14 days

The study consisted of exposing six male albino rats to a flowing stream of air with sBA vapors. The stream was prepared by proportioning pumps and nominal concentrations (not confirmed by analytical methods) were recorded. The study end point was the concentration yielding a fractional

mortality among six rats within 14 days.

Result : 8,000 ppm sBA was lethal to 1 of 6 rats within 14 days

Conclusion: One of six exposed rats died within 14 days.

Reliability : (2) valid with restrictions

2- Reliable with restrictions. Pre-GLP study documented, meets generally

accepted scientific principles, acceptable for assessment.

10.01.2002 (65)

Type : LC50

Date

Value : = 10000 ppm

Species : rat

Strain : Sprague-Dawley

Sex : female Number of animals : 5

Vehicle : other: None

Doses

Exposure time : 7 hour(s)

Method : other: Experimental (Non-regulatory)

Year : 1989 GLP : yes

Test substance : as prescribed by 1.1 - 1.4

Remark : Study Type: Acute Inhalation (Pilot for Teratology study)

Purity: >or= 99 %

Route of Administration: Inhalation Observation Period: Unknown

The study consisted of exposing five female rats to a flowing stream of air with sBA vapors. The stream was prepared by proportioning pumps. The study end point was the concentration yielding a fractional mortality.

Result : 10,000 ppm sBA was lethal to 5 of 5 rats

Conclusion: Five of five exposed female rats died following a single 7 hour exposure.

Reliability : (2) valid with restrictions

2 - Reliable. Pilot study with acceptable restrictions.

10.01.2002 (69)

Type : LC50

Value

Species: mouseStrain: no dataSex: no data

Number of animals : 2

Vehicle : other: None

Doses

Exposure time

Method : other: Experimental (Non-regulatory)

Year : 1938 GLP : no

Test substance : as prescribed by 1.1 - 1.4

Remark: 1650 ppm: no signs of intoxication after 420 minutes.

3300 ppm: ataxia in 51-100 minutes, prostration in 120-180 minutes,

narcosis in 3000 minutes, no deaths

19,800 ppm: ataxia in 7-8 minutes, prostration in 12-20 minutes, narcosis in

40 minutes, no deaths

Route of Administration: Inhalation

Doses/time: 1650, 3300, 19800 ppm/decreasing lengths of time

Post Dose Observation Period: Various

Result : Not Defined

Conclusion: Doses from 3,300 to 19,800 ppm induced narcosis but not death.

Reliability : (4) not assignable

4 - Documentation insufficient for assessment. Non-guideline pre-GLP

study.

10.01.2002 (85)

5.1.3 ACUTE DERMAL TOXICITY

Type : other: Limit
Value : > 2000 mg/kg bw

Species : rat

Date

Strain : Fischer 344
Sex : male/female

Number of animals : 10

Vehicle : other: None

Doses

Method : other: Experimental (Non-regulatory)

Year : 1986 **GLP** : yes

Test substance : as prescribed by 1.1 - 1.4

Remark: Fischer 344 rats were in the age range of 55-65 days when purchased and

were 9 -11 weeks old when the study commenced. The day before dosing the animals, approximately 60% of the dorsal hair was closely shorn with fine electric clippers, and the rats and the rats were weighed on the day of dosing. Varying the volume dispensed from the syringe altered doses. Application was made at room temperature and the test material was covered with aluminum foil and held in place by a double over-wrap of waterproof adhesive tape. The rats were individually housed for the next 24 hours. Food was withheld, but water was available ad libitum. At the end of the 24-hour period, the tape and foil were removed and the skin was washed with warm dilute detergent solution and then dried. The animals were observed for signs of toxicity for 14 days after dosing. Body weights were recorded on Days 1, 7 and 14. None of the rats died. There were no overt signs of toxicity and all rats gained weight relative to their day 1body

weight.

Route of Administration: Dermal

Doses/time: 2000 mg/kg single application / 24-hour occlusive patch

Post Dose Observation Period: Daily for 14 Days

Purity: 99.5% > 2000 mg/kg

Result : > 2000 mg/kg

Conclusion : The acute percutaneous LD50 of undiluted sBA in rats was > 2000 mg/kg.

Reliability : (1) valid without restriction

1 - Reliable without Restrictions. No circumstances occurred that would

have affected the quality or integrity of the data.

10.01.2002 (73)

5.1.4 ACUTE TOXICITY, OTHER ROUTES

Type : LD50

Value : = 800 mg/kg bw

Species : mouse

Strain

Sex :
Number of animals :
Vehicle :
Doses :

Route of admin. : i.p. Exposure time :

Method : other: not specified

Year : 1978 GLP : no data

Test substance: as prescribed by 1.1 - 1.4

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

21.10.1992

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5.2.1 SKIN IRRITATION

Species : rabbit

Concentration : Exposure : Exposure time : Number of animals : Vehicle : PDII :

Result : not irritating
Classification : not irritating
Method : other: not specified

Year : 1954 GLP : no data

Test substance: as prescribed by 1.1 - 1.4

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

21.10.1992 (33)

Species : rabbit

Concentration

Exposure : Occlusive **Exposure time** : 4 hour(s)

Number of animals : Vehicle : PDII :

Result : not irritating
Classification : not irritating

Method : OECD Guide-line 404 "Acute Dermal Irritation/Corrosion"

Year : 1986 **GLP** : yes

Test substance: as prescribed by 1.1 - 1.4

Remark: New Zealand White rabbits, 3 - 6 months of age, were acclimated to the

laboratory conditions for at least 2 weeks prior to experiment initiation. The hair of the dorsal back between the shoulders and hindquarters of three male and 3 female rabbits, 4 - 9 months of age at dosing, was closely shorn with fine electric clippers. A test site was selected and a 2 cm x 2 cm lint patch with 0.5 ml of sBA was applied to it. The patch and surrounding skin were covered by a single layer of gauze and held in place with an elastic adhesive bandage. The wrapping and patch were removed after 4 hours and the skin was not washed. The site was examined and scored for erythema and edema on a graded scale 0 - 4. Observations were made 30 minutes after the removal of the patch and at 24, 48, and 72 hours and 7 days following dosing. The mean scores for each rabbit at each

observation time were calculated. There were no skin reactions following the application of sBA to rabbit skin for 4 hours. The test material is

therefore not a skin irritant in rabbits.

Purity: 99.5%

Sex: Male and Female

Vehicle: None

Route of Administration: Dermal

Doses: 0.5 ml

Doses/Time: Single application

Post Dose Observation Period: After removal of patch: 0.5 hours; 24, 48,

72 hours and Day 7 after dosing.

Result : PII = 0

Conclusion: sBA caused no skin reaction and is not a skin irritant in rabbits.

Reliability : (1) valid without restriction

1 - Reliable without Restrictions. No circumstances occurred that would

5. Toxicity Id 78-92-2

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have affected the quality or integrity of the data

10.01.2002 (73)

5.2.2 EYE IRRITATION

Species : rabbit

Concentration :
Dose :
Exposure time :
Comment :
Number of animals : 5
Vehicle :

Result : irritating Classification : irritating

Method : other: Experimental (Non-regulatory)

Year : 1954 GLP : no

Test substance : as prescribed by 1.1 - 1.4

Remark: Eye injury in rabbits records the degree of corneal narcosis from various

volumes and concentrations of chemical, as detailed in Carpenter and Smyth (1946). Grade 1 indicates at most a very small area of narcosis resulting from 0.5 ml of undiluted chemical in the eye; Grade 5 indicates a so-called severe burn from 0.005 ml, and Grade 10 indicates a severe burn

from 0.5 ml of a 1% solution in water or propylene glycol.

Sex: Male Vehicle: None

Route of Administration: Ocular application into the conjunctival sac of one

eye

Control: Untreated eye

Dose: Single application of undiluted sBA

Volume: 0.02 and 0.1 ml

Post Dose Observation Period: 24 Hours

Result: Irritating (score 4/10); Severe injury from 0.1 ml, minor from 0.02 ml; Grade

4 eye injury.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Conclusion : sBA caused moderate damage and/or corneal opacity in rabbits.

Reliability : (2) valid with restrictions

2 - Reliable with restrictions: data are from a collection of data.

10.01.2002 (17) (82)

Species : rabbit

Concentration :

Dose :

Exposure time :

Comment :

Number of animals :

Number of animals : 6 Vehicle :

Result : corrosive

Classification

Method : OECD Guide-line 405 "Acute Eye Irritation/Corrosion"

Year : 1986 **GLP** : yes

Test substance: as prescribed by 1.1 - 1.4

Remark: New Zealand White rabbits, 3 - 6 months of age, were acclimated to the

laboratory conditions for at least 2 weeks prior to experiment initiation. Three male and 3 female rabbits were 4 - 9 months of age at dosing. The day before testing, the eyes were carefully examined for any damage and

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any animals showing damage were replaced. A single dose of 0.1 ml of sBA was placed into the lower conjunctival sac of one eye and the lids held together for a few seconds to prevent loss of material. The eyes were not washed. The reactions of the animal were observed and the initial pain responses graded on a 1 - 6 scale, no pain to very sever initial pain. Visual assessments were made 1, 5-7, 24, 28, and 72 hours and 7 Days postdosing. Irritancy was scored for the cornea, iris, and conjunctiva using standard scores. Any corneal damage visualization was aided by instillation of one drop of 2% fluorescein solution. The degree of irritation was classified using a scheme based on the OECD Data Interpretation Guide (1984). The instillation of undiluted sBA resulted in moderate initial pain. The conjunctival redness, chemosis and discharge, corneal opacity and damage to the iris were assessed and mean scores calculated. All rabbits had moderate conjunctival inflammation with some discharge within 1 hour of dosing. The swelling and discharge largely cleared by four hours but the redness persisted in 3 rabbits for 7 days. These 3 animals, and another, had impaired iritic response and/or slight corneal opacity between 24 and 72 hours postdosing. The effect had cleared by 7 days in 3 rabbits. The severely affected rabbit had a completely opaque cornea and no iritic response. It was humanely terminated since recovery was deemed not possible. The remaining rabbits were retained and by day 14 all ocular effects had cleared. In view of the responses, sBA is classified as corrosive to rabbit eyes (OECD, 1984).

Study Type: Ocular Irritation (Draize type)

Purity: 99.5%

Sex: Male and Female

Vehicle: None

Route of Administration: Ocular application into the conjunctival sac of one

eve

Control: Untreated Eye

Dose: Single application of neat material

Volume: 0.1 ml

Post Dose Observation Period: 1, 5-7, 24, 28, and 72 hours and 7 Days

postdosing

Result : Corrosive

Conclusion: sBA caused moderate conjunctival inflammation in all six rabbits with slight

transitory iritic damage and/or corneal opacity in three rabbits. One rabbit developed intense, extensive corneal opacity and complete loss of iritic response. The test material was there for corrosive to the rabbit eye. On administration of sBA, there was a moderate initial pain response.

Reliability : (1) valid without restriction

1 - Reliable without Restrictions. No circumstances occurred that would

have affected the quality or integrity of the data.

10.01.2002 (73)

5.3 SENSITIZATION

Type : Freund's complete adjuvant test

Species : guinea pig

Number of animals : 20

Vehicle: other: Corn oilResult: not sensitizingClassification: not sensitizing

Method : other: Magnusson and Kligman; 1969

Year : 1986 GLP : yes

Test substance: as prescribed by 1.1 - 1.4

Remark: Guinea pigs in the weight range of 300-370 grams were purchased from a

commercial supplier and acclimated to the laboratory for at least 2 weeks.

Date

The skin sensitization potential of sBA was assessed using the guinea pig maximization test of Magnusson and Kligman. The test was accomplished in 2 stages: rangefinding and definitive test.

Rangefinding:

The purpose of the rangefinding studies was to determine the concentrations of test material to be used for intradermal induction, topical induction and topical challenge. Two males and two female guinea pigs were closely shorn in the shoulder region using electric clippers followed by an electric razor and 0.1 ml of several dilutions (0.05, 0.1, 0.5 and 1.0% [m/v] in corn oil) of sBA injected intradermally each side of the midline. The animals were observed over the next few days to determine the maximum concentration that could be tolerated without causing untoward toxicity. Three further groups of two males and two females had their flanks closely shorn and 0.3 ml of several dilutions (25%, 50%, and 75% [m/v] in corn oil) of sBA were applied to a 4 cm x 4 cm Whatman Number 3 filter paper patches. The patches were placed on the flanks and held in place with a "Sleek" adhesive tape patch, then covered with a "Poroplast" elastic adhesive bandage for 24 hours. The bandages were removed and the animals were examined for signs of irritation and scored using a 4-point scale (0, 1, 2, and 3). The concentration used for topical induction was the one that scored 1 for irritation and the concentration for challenge was the one that was 0, a non-irritant.

Definitive:

This test was conducted using a group of ten male and ten female guinea pigs together with a control group of five males and five females. The test consisted of two stages: a) induction by intradermal injection and topical application, and, 2) topical challenge. The following concentrations were selected for the definitive study: Intradermal induction, 0.1% (m/v) in corn oil; Topical induction (one week following intradermal challenge), 50% (m/v) in corn oil and 25% (m/v) in corn oil. Topical challenge was carried out two weeks following the topical induction by applying 0.1 ml of the diluted sBA to a semi-occluded challenge patch and removing it 24 hours later. The erythema resulting from the topical challenge was scored on a 4-point scale (0, 1, 2, and 3) immediately on removal of the challenge patches and 24 and 48 hours latter. None of the twenty test animals showed any positive response at either 24 or 48 hours after the removal of the challenge patches. It was concluded that sBA was not a dermal sensitizer in guinea pigs.

Purity: 99.5%

Sex: Male and Female

Route of Administration: Intradermal and topical induction; topical challenge

Doses:

Intradermal Induction:

Test Doses:

2 injections (0.1 ml) of Freund's Complete Adjuvant (FCA)

2 injections (0.1 ml) of sBA in corn oil

2 injections (0.1 ml) of sBA in 50:50 FCA:corn oil

Control Doses:

2 injections (0.1 ml) of Freund's Complete Adjuvant (FCA)

2 injections (0.1 ml) of corn oil

2 injections (0.1 ml) of 50:50 FCA:corn oil

Topical Induction:

Application of 0.3 ml 50% sBA (m/v) in corn oil and, 25% sBA (m/v) in corn oil covered for 48 hours.

Topical Challenge:

Application 0.1 ml of diluted sBA to a semi-occluded challenge patch and

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removing it 24 hours later.

Post Dose Observation Period: Immediately, 24 and 48 hours following

challenge

Result : sBA is not a dermal sensitizer in guinea pigs.

Conclusion : In the guinea pig maximization test of Magnusson and Kligman, sBA did

not induce positive responses in any of the 20 test animals at 24 or 48 hours after removal of the challenge patches. sBA is therefore not a

dermal sensitizer in guinea pigs.

Reliability : (1) valid without restriction

1 - Reliable without Restrictions. No circumstances occurred that would

have affected the quality or integrity of the data.

11.01.2002 (73)

Type : Guinea pig maximization test

Species : guinea pig

Number of animals : 20

Vehicle: other: Paraffin oilResult: not sensitizingClassification: not sensitizing

Method : other: OECD Guide-line 406 and EC92/69/EEC, B6

Year : 1992 GLP : yes Test substance : other TS

Remark : Age: ~ 3 months

Sex: Male and Female

Weight (initial): Males 335 ± 17 g; Females 332 ± 19 g

Doses:

Intradermal Induction:

Test Doses:

Injections: sBA in paraffin oil (5% w/w) sBA in FCA (Freund's Complete Adjuvant)

Control Doses: Injections: Paraffin oil

Freund's Complete Adjuvant (FCA)

Topical Induction:

Application: sBA (undiluted) covered for 48 hours

Topical Challenge:

Application: sBA (undiluted) covered for 24 hours

Post Dose Observation Period: Skin reactions were evaluated at

approximately 24 and 48 hours.

Clinical Signs: None

Rechallenge: None

Thirty guinea pigs were allocated to two groups: a control group 1 (five males and five females) and a treated group 2 (ten males and ten females). Day 1: Intradermal injections of Freund's complete adjuvant mixed with the test substance (treated group) or the vehicle (control group) were performed in the dorsal region between the shoulders. Day 7: The same dorsal region between the shoulders received a topical application of sodium lauryl sulfate in Vaseline (10% w/w) in order to induce local irritation.

Day 8: The test site was treated by topical application of the test substance (treated group) or the vehicle (control group) and was covered by an occlusive dressing for 48 hours.

Day 22: After a rest period of 12 days, all animals of the treated and control

Date

groups were challenged by a topical application of the test substance to the right flank. The left flank served as control and received the vehicle only. Test substance and vehicle were maintained under an occlusive dressing for 24 hours. Skin reactions were evaluated approximately 24 and 48 hours later.

At the end of the study, animals were killed without examination of internal organs. No skin samples were taken from the challenge application sites.

The sensitivity of the guinea pigs was checked with a positive sensitizer: 2,4-dinitro chlorobenzene (DNCB). During the induction period, the test substance was applied at 0.1 % (w/w) (day 1) and 1 % (w/w) (day 8). For the challenge application, the DNCB was applied to the right flank at a

concentration of 0.5% (w/w).

Result : sBA is not a dermal sensitizer in guinea pigs. The sensitivity of the guinea

pigs was satisfactory since 50% of the animals showed a positive reaction with positive control sensitizer, DNCB. Scores in control and treated groups

after challenge with sBA were 0 at both 24 and 48 hours.

Source : Centre d'Elevage Lebeau, Gambais, France

Test substance: sec-Butanol (sBA) 99.87%; ATOFINA Chemicals, Inc,

Conclusion: In the guinea pig maximization test of Magnusson and Kligman, sBA is not

a dermal sensitizer.

Reliability : (1) valid without restriction

1 - Reliable without Restrictions.

11.01.2002 (32)

5.4 REPEATED DOSE TOXICITY

Type : rat Sex : male

Strain : Sprague-Dawley
Route of admin. : inhalation
Exposure period : 3-5 days
Frequency of treatm. : 6 hours/day

Post exposure period

Doses : 2000 ppm (3 Days) and 500 ppm (5 Days)

Control group :

Method : other: Experimental -- Study of changes in cytochrome P-450 enzyme

system of rat liver, kidney and lung

Year : 1985 GLP : no data

Test substance: as prescribed by 1.1 - 1.4

Remark : Purity: > or = 99%

Number/Sex/Dose: 5

Vehicle: None

Result : sBA caused increases in cytochrome P-450 concentrations in the kidneys

(47% rise; 500 ppm for 5 days) and liver (33% rise; 2000 ppm for 3 days).

Slight decreases in lung cytochrome P-450 were observed.

Conclusion : sBA (probably through its metabolite methyl-ethyl-ketone) was a potent

inducer of P-450 in the kidney and liver.

Reliability : (2) valid with restrictions

2- Reliable study with restrictions. Study well documented, meets generally accepted scientific principles, acceptable for assessment.

11.01.2002 (2)

Type : Species : rat

Sex : male/female

Date

Strain : Fischer 344
Route of admin. : inhalation
Exposure period : 90 days

Frequency of treatm. : 6 hours/day; 5 days/week

Post exposure period

Doses : 0, 1250, 2500, or 5000 ppm - Vapor

Control group : yes

NOAEL : = 2500 ppm **LOAEL** : = 5000 ppm

Method : other: Comparable to OECD Guideline 413; 90-Day Subchronic Toxicity

Year : 1981 GLP : yes Test substance : other TS

Remark: An extensive pathologic investigation was conducted and no lesions were

found that could be attributed to MEK exposure. There was no indication that repeated exposure to relatively high levels of MEK had any effect on reproductive tissues. The examined tissues included testes, epididymides,

seminal vesicles, vagina, cervix, uterus, oviducts, and ovaries. Methyl ethyl ketone is metabolically interchangeable with 2-butanol. Approximately 97% of a 2-butanol dose is

oxidized to methyl ethyl ketone.

Purity: > 99.5%

Number/Sex/Dose: 15 (10/sex - principals - for routine pathology and

5/sex - dedicated - for special neuropathology)

Vehicle: None

Result: The 90-day exposures had no adverse effect on the clinical health or

growth of male or female rats except for a depression of mean body weight in the 5000-ppm exposure group. No animals died during the study. No signs of nasal irritation were observed during the study. There were no treatment-related effects in food consumption or ophthalmologic studies in any of the rats exposed to MEK vapors. The 5000-ppm animals had a slight but significant increase in liver weight, liver weight/body weight ratio and liver weight/brain weight ratio at the time of necropsy. Serum glutamic-pyruvic transaminase (SGPT) activity in the 2500-ppm female rats was elevated while the 5000-ppm female rats exhibited significantly decreased SGPT activity. In addition, alkaline phosphatase, potassium, and glucose

values for the 5000-ppm female rats were increased. Special

neuropathological and routine pathological studies did not reveal any

lesions that could be attributed to MEK exposure.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Test substance : Methyl Ethyl Ketone (MEK)

Conclusion: MEK has a low order of toxicity based on the 90-day repeated-dose

exposures. The NOAEL is 2500 ppm.

Reliability : (1) valid without restriction

1 - Reliable study with restrictions. Study well documented, meets generally accepted scientific principles, acceptable for assessment.

14.01.2002 (19) (89)

5.5 GENETIC TOXICITY 'IN VITRO'

Type : Ames test System of testing : Bacterial

Test concentration : 0.01, 0.1, 0.5, 1.0, and 5.0 mg/ml

Cycotoxic concentr.

Metabolic activation: with and withoutResult: negative

Method : OECD Guide-line 471

Year : 1985

GLP : yes

Test substance: as prescribed by 1.1 - 1.4

Remark : Purity: 99.5%

Strain: S. typhimurium / TA98, TA100, TA1535, TA1537, TA1538

Species/Cell Type: Rat Liver (S9 fraction)

Vehicle: DMSO

sBA did not induce reverse gene mutation in bacteria.

EXXON CHEMICAL, Limited Fareham, Hampshire

Source : EXXON CHEMICAL, Limited Fareham, Hampshire EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Test condition : Control plates were set up with solvent alone and with an appropriate

known positive control compound. All tests were carried out in triplicate. Two replicate assays were carried on different days in order to confirm the reproducibility of the results. The S9 fractions were prepared from Aroclor-induced rats. Initially, a range of test concentrations of test material was tested and a second experiment was then performed based on the results taking into account the effect on cell viability and any possible positive increases in mitotic gene conversion. Control plates were set up with solvent alone and with the positive control compounds 4-nitroquinoline-N-

oxide and cyclophosphamide.

Conclusion : Not Mutagenic

Reliability : (1) valid without restriction

1- Reliable without restriction

11.01.2002 (15)

Type : Ames test System of testing : Bacterial

Test concentration : 0.01, 0.1, 0.5, 1.0, and 5.0 mg/ml

Cycotoxic concentr.

Test condition

Metabolic activation : with and without

Result : negative

Method : OECD Guide-line 472

Year : 1985 **GLP** : yes

Test substance: as prescribed by 1.1 - 1.4

Remark : Purity: 99.5%

Strain: E. coli WP2 uvr A

Species/Cell Type: Rat Liver (S9 fraction)

Vehicle: DMSO

sBA did not induce reverse gene mutation in bacteria.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA) Control plates were set up with solvent alone and with an appropriate

known positive control compound. All tests were carried out in triplicate. Two replicate assays were carried on different days in order to confirm the reproducibility of the results. The S9 fractions were prepared from Aroclor-induced rats. Initially, a range of test concentrations of test material was tested and a second experiment was then performed based on the results taking into account the effect on cell viability and any possible positive increases in mitotic gene conversion. Control plates were set up with solvent alone and with the positive control compounds 4-nitroquinoline-N-

oxide and cyclophosphamide.

Conclusion : Not Mutagenic

Reliability : (1) valid without restriction

1- Reliable without restriction.

14.01.2002 (15)

Type : other: Yeast mitotic gene conversion

System of testing : Saccharomyces cerevisiae Test concentration : Maximum conc. 5 mg/ml

Cycotoxic concentr. :

59 / 76

Metabolic activation : with and without

Result : negative

Method : OECD Guide-line 481

Year : 1988 **GLP** : yes

Test condition

Test substance: as prescribed by 1.1 - 1.4

Remark : Purity: 99.5%

Strain: Saccharomyces cerevisiae Species/Cell Type: Rat Liver (S9 fraction)

Vehicle: DMSO

sBA did not induce mitotic gene conversion in yeast.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA) : Yeast cells were grown in log phase, washed and re-suspended in 2/5

strength YEPD broth at a concentration of 10 X 106cells/ml. The suspension was then divided into 1.9 ml amounts in 30-ml universal containers and 0.1 ml of the test compound solution was added (-S9). For the experiments with metabolic activation (+S9), 0.1 ml of the compound was added to 1.6 ml of yeast suspension, together with 0.3 ml of S9 mix. The cultures were incubated with shaking, with shaking, at 30°C for 18 hours. Aliquots were then plated onto the appropriate culture media for the selection of mitotic gene convertants and cells surviving the treatment.

Conclusion : Not mutagenic

Reliability : (1) valid without restriction

1- Reliable without restriction

14.01.2002 (14)

Type : other: chromosome abberration assay

System of testing : Chinese hamster ovary cells
Test concentration : maximum conc. 5000 ug/ml

Cycotoxic concentr.

Test condition

Metabolic activation: with and without

Result : negative

Method : OECD Guide-line 473

Year : 1988 **GLP** : yes

Test substance: as prescribed by 1.1 - 1.4

Remark : Purity: 99.5%

Species/Cell Type: Chinese Hamster Ovary Cells

Vehicle: DMSO

sBA did not induce chromosome damage in CHO mammalian cells.

Result : Negative - No increase in chromosomal aberrations
Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

: Two separate cytotoxicity assays were performed: (i) Monolayer CHO cultures were prepared in multi-well tissue culture travs. The cultures were

incubated at 37°C for 24 hours to commence active growth before treatment with the test material. Assays were performed both in the presence and in the absence of S9 mix using either a 5-hour exposure

(+S9) or a 24-hour exposure (-S9). Twenty-four hours after initial

treatment, cultures were stained with Giemsa and the growth inhibition was noted. (ii) CHO cultures were prepared in 25 cm2 flasks, and after 24 hours the cells were exposed to sBA both in the presence and in the absence of S9 mix. The number of cell in each flask was counted 24 hours later. The compound concentrations selected for the chromosome assays

were 1, 0.5, and 0.25 times the GI50 level.

CHO Chromosome Assay: Cultured CHO cells were grown in 80-cm2 flasks for 24 hours before treatment. Treatment periods were 5hours in the presence of S9 mix and 24 hours in the absence of S9 mix. Positive control

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cultures were run in parallel [ethyl methanesulphonate (-S9) and cyclophosphamide (+S9). Colcemid was added to all cultures 22 hours after treatment. After a further 2 hours, the cells were trypsinized, resuspended in hypotonic solution and then fixative, before spotting onto slides. Cell preparations were then stained with Giemsa. The slides were randomly coded and 100 cells from each culture were analyzed

microscopically. Mitotic index estimations were also made. Not Mutagenic

Reliability (1) valid without restriction

1- Reliable without restriction

14.01.2002 (14)

Type Ames test System of testing **Bacterial**

Test concentration 100-500-1000-5000-10000 ug/plate

Cycotoxic concentr.

Conclusion

Metabolic activation with and without

Result negative

Method OECD Guide-line 471

Year 1988 **GLP** yes

Test substance as prescribed by 1.1 - 1.4

sBA did not induce reverse gene mutation in bacteria. Remark

Strain: Salmonella typhimurium / TA98, TA100, TA1535, TA1537, TA1538

Species/Cell Type: Rat Liver (S9 fraction)

Vehicle: DMSO

Source EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Test condition Control plates were set up with solvent alone and with an appropriate

> known positive control compound. All tests were carried out in triplicate. Two replicate assays were carried out on different days in order to confirm the reproducibility of the results. The S9 fractions were prepared from Aroclor -induced rats. Initially, a range of test concentrations of test material was tested and a second experiment was then performed based on the results taking into account the effect on cell viability and any

possible positive increases in mitotic gene conversion. Control plates were set up with solvent alone and with the positive control compounds 4-

nitroquinoline-N-oxide and cyclophosphamide.

Conclusion : Not Mutagenic

: (1) valid without restriction Reliability

1- Reliable without restriction

11.01.2002 (31)

5.6 GENETIC TOXICITY 'IN VIVO'

Type other: In Vivo Mammalian Bone Marrow Erythrocyte Micronucleus Assay

Intraperitoneal Dosing Method

Species mouse male/female Sex

Strain CD-1

Route of admin. other: Intraperitoneal injection

12, 24 and 48 hours Exposure period 1.96 ml/kg Single injection Doses

Result

other: OECD 474 equivalent Method

Year 1988 **GLP** no data Test substance other TS

Remark : Not mutagenic

Purity: 99.9% Number: 5/sex/dose

Dose/Time: 1.96 ml/kg Single injection (dose equal to the MEK LD20 reported as ml test article / kg body weight when administered in a total

volume of 10 ml test article-vehicle mixture / kg body weight).

Vehicle: Corn Oil (10 ml/kg)

Positive Control: 0.25 mg/kg Triethylene melamine (TEM)

The test substance and the vehicle were administered as a single dose by intraperitoneal injection. The vehicle was dosed at a volume equal to the test substance volume. The positive control was administered as a single dose at a volume equal to the test substance volume. Animals from the appropriate groups were sacrificed at approximately 12, 24 and 48 hours. Animals dosed with triethylene melamine were sacrificed at 24 hours only. Immediately following sacrifice, the femur was exposed and the bone marrow was aspirated into a syringe containing fetal calf serum. The cells were washed, centrifuged, and resuspended. Slide smears of the bone marrow were made for each animal and stained with May-Gruenwald-Giemsa stain. Coded slides were then evaluated for presence of micronuclei (1000 polychromatic erythrocytes/animal were evaluated). A 1-way analysis of variance and Duncan's multiple range test (p = 0.05) were used to assess the statistical significance of any observed effects.

Result: None of the dose groups were statistically different from the vehicle control.

The positive control (0.25 mg/kg TEM) induced a statistically significant increase in the mean number of micronucleated polychromatic erythrocytes (p = 0.01) which indicates that the positive control was clastogenic and the test system responded in an appropriate manner. Vehicle carrier control values for the mean percent of polychromatic erythrocytes and for the mean percent of micronucleated polychromatic erythrocytes responded in

an appropriate manner.

Test substance: Methyl Ethyl Ketone (MEK) CAS No. 78-93-3

Conclusion : MEK did not induce a statistically significant increase in the mean number

of micronucleated polychromatic erythrocytes in the bone marrow of CD-1 mice. Therefore, it is not considered mutagenic under the conditions of this

assay.

Reliability : (1) valid without restriction

1 - Reliable without Restrictions. No circumstances occurred that would

have affected the quality or integrity of the data.

29.01.2002 (70)

Type : other: Rat Bone Marrow

Species : rat
Sex : no data
Strain : no data

Route of admin. : other: intragastric

Exposure period :
Doses :
Result :

Method : other: Micronucleus Aberration

Year : 1988 GLP : no data

Test substance: as prescribed by 1.1 - 1.4

Remark : Single intragastric administration of monohydric alcohols at equitoxic doses

(1/5 LD50) affected the chromosomal appearance of bone marrow.

System of Testing: Rat Bone Marrow Metabolic Activation: unknown

Species/Cell Type: Rat

Concentrations Tested: 1/5 of the LD50 by intragastric route

Vehicle: unknown

Result: Increased numbers of polyploid cells, cells with chromosome gaps, and

cells with chromosomal aberrations.

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Conclusion: unknown

Reliability : (4) not assignable

4 - Incomplete translation from Russian and insufficient for assessment

11.01.2002 (5)

5.7 CARCINOGENICITY

Remark: No studies of 2-butanol for carcinogenic activity have been

reported; however, based on results of mutagenicity assays it is expected to have a low potential for carcinogenicity.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

21.10.1992

5.8.1 TOXICITY TO FERTILITY

Type : other: two generation study with teratology screen

Species : rat

Sex : male/female
Strain : Wistar
Route of admin. : drinking water

Exposure period :

Frequency of treatm. : Daily - ad libitum

Premating exposure period

Male : 8 weeks
Female : 8 weeks
t : 2 Generations

Duration of test : 2 Generat

No. of generation

studies

Doses : F0 Generation: 0, 0.3, 1.0, or 3.0% solutions; F1 Generation: 0, 0.3, 1.0, or

2.0%

Control group : yes NOAEL F1 offspring : = 1 % other: NOAEL : = 1 %

Maternal

Method : other: Comparable to an OECD 416 guideline study.

Year : 1975 **GLP** : no

Test substance: as prescribed by 1.1 - 1.4

Remark : Purity: > or = 99%

Number/Sex/Dose: 30

Doses/Concentration:

F0 Generation: 0, 0.3, 1.0, or 3.0% solutions (0, 538, 1644, and 5089 mg/kg-day for males and 0, 594, 1771, and 4571 mg/kg-day for females) F1 Generation: 0, 0.3, 1.0, or 2.0% (2.0% calculated to be equivalent to

3384 mg/kg-day for males and 3122 mg/kg-day for females)

Vehicle: Drinking Water

sec-Butanol was initially administered to the F0 generation at concentrations of 0, 0.3, 1.0, and 3.0% in the drinking water. Due to toxicity, the high level was reduced to 2.0% during the second-generation study (F1). The F1 generation animals (30/sex/group) were reared to maturity (up to week 12), mated to produce a F2 generation, then sacrificed for organ weights, and gross and microscopic pathological evaluations

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(10/sex/group). Hematological, biochemical, and urinary examinations were conducted terminally on the F1 rats. A series of mild changes in the kidney (non-reactive tubular degeneration, tubular casts, foci of tubular regeneration, microcysts) were observed animals treated at 2.0% sBA. The authors concluded that these findings were not a result of direct toxicity and did not have clear pathologic significance. Rather they were non-specific effects due to increased renal work load, possibly from increased urine volume and pressure at the high dose of sBA (Cox et al. 1975). No other findings of note were seen. The no-effect level for the study was 1.0% (estimated to be 1500 mg/kg/day by the authors and 1771

mg/kg/day by EPA/IRIS).

Result Maternal NOEL: 1% (~1500 mg/kg/day)

> Maternal NOAEL: 1771 mg/kg/day Pup NOEL: 1% (~1500 mg/kg/day) Pup NOAEL: 1771 mg/kg/day

EXXON CHEMICAL, Limited Fareham, Hampshire Source

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

Conclusion sBA is not a reproductive hazard.

(2) valid with restrictions Reliability

> 2 - Reliable study with restrictions. No circumstances occurred that would have affected the quality or integrity of the data. Comparable to a guideline study and test procedures were in accordance with generally accepted

scientific standards and described in sufficient detail.

11.01.2002 (22)(40)

5.8.2 DEVELOPMENTAL TOXICITY/TERATOGENICITY

Species : rat Sex female

Strain Sprague-Dawley Route of admin. inhalation

Gestation Days 1-19 Exposure period

Frequency of treatm. 7 hr/day **Duration of test** 20 days

3500, 5000, 7000 ppm **Doses**

Control group yes

NOAEL maternal tox. = 3500 ppm : > 7000 - ppm NOAEL teratogen.

Method other: Comparable to an OECD 414 guideline study

Year 1989 **GLP** ves

Test substance as prescribed by 1.1 - 1.4

Method For the maternal data, multivariate analysis (with baseline as covariant)

> was used for weight comparisons across groups. The group differences in food and water intake were analyzed by multivariate analysis of variance. A Kruskal-Wallis test was used for group comparisons of corpora lutea per animal. For the fetal data, analysis of covariance was used to compare fetal weights across groups and sex. Group comparisons of the variables including litter size, percentage alive/litter, percentage normal/litter, and percentage females/litter were made using Kruskal-Wallis test. For the variables including skeletal malformations, skeletal variations, visceral malformations, visceral variations, external malformations, and non-normal fetuses, the number of litters with one or more of the variables of interest was compared between groups using Fisher's exact test. The results of the test were adjusted for multiple comparisons, when appropriate, using the Bonferroni technique. A probability of p = < 0.05 was required for

significance.

Remark Groups of 15-16 rats were exposed by inhalation to 0, 3,500, 5,000 or

7,000 ppm sBA, 7 hours/day on days 1-19 of gestation; the dams were

sacrificed on day 20. At 7,000 ppm, narcosis was observed in all animals. At 5000 ppm, the dams were partially narcotized with locomotion activity impaired. Maternal weight gain and food consumption were significantly reduced in all dose groups. No data collected on maternal organ weights, or gross or microscopic lesions. The number of live fetuses was significantly reduced and resorptions were increased in the high exposure group only. Fetal body weights were significantly reduced in the mid- and

group only. Fetal body weights were significantly reduced in the mid- and high dose groups. There was no evidence of teratogenic effects in this study, and there was also no evidence of selective developmental toxicity. The no-effect levels were < 3,500 ppm for maternal toxicity and 3,500 ppm

for developmental toxicity. Purity: > or = 99%

Number/Sex/Dose: 15-16 Mated Females

Vehicle: None

Result : Maternal NOEL: < 3500 ppm

Maternal NOAEL: 3500 ppm Pup NOEL: 3500 ppm Pup NOAEL: > 7000 ppm

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)sBA is not a teratogen. There was no evidence of teratogenic events nor

was there evidence of selective developmental toxicity.

Reliability : (2) valid with restrictions

2- Reliable study with restrictions. Study well documented, meets generally accepted scientific principles, acceptable for assessment.

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Species: ratSex: femaleStrain: Wistar

Conclusion

Route of admin. : drinking water

Exposure period : 8 Weeks premating (Males and Females) and during gestation (Females)

Frequency of treatm. : Daily - ad libitum

Duration of test : 20 days

Doses : 0, 0.3, 1.0, or 2.0% solutions

Control group : yes

Method : other: Comparable to an OECD 421 guideline study.

Year : 1975 **GLP** : no

Test substance: as prescribed by 1.1 - 1.4

Remark : Purity: > or = 99%

Doses/Concentration: F0 Generation - second breeding.

 $0,\, 0.3,\, 1.0,\, or\, 2.0\% \,\, solutions \,\, (0,\, 538,\, 1644,\, and\, 3384 \,\, mg/kg-day \,\, for\, males$

and 0, 594, 1771, and 3122 mg/kg-day for females)

Vehicle: Drinking Water Number/Sex/Dose: 30

TERATOLOGY SCREEN RESULT: The F0 rats were mated to obtain a second series of pregnant dams destined to provide teratologic evaluation of the treatments. Pregnancy rates and survival of these females were unaffected. All findings sBA at both 0.3 and 1.0% in the drinking water were negative with respect to signs of toxicity in terms of both growth and reproductive efficiency The body weights of the dams were not depressed. Examination of the uterine contents on the 20th day of gestation revealed that sBA was somewhat fetotoxic at the 2.0% dosage level, as shown by the decreased pup weights (3.74 g vs. 4.14 g in controls). However, that this is a minimal response is shown by the fact that none of the other parameters in the reproductive toxicity phase of this study (nidation, early or late fetal deaths) were affected. The 2.0% group showed apparent increases in missing sternebrae, wavy ribs, and incomplete vertebra ossification when compared with both the 0.3 and 1.0% groups. However, because the incidences for these findings in the control group were

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comparable, these effects could not be determined to be compoundrelated. The skeletal abnormalities seen in the sBA groups were consistent in type and frequency with the spontaneous incidence observed in this rat colony. There were no significant soft tissue findings in the 2% treated

group.

Result : Maternal NOEL: 1% (1771 mg/kg/day)

Maternal NOAEL: 1% (1771 mg/kg/day)

Pup NOEL: 1% (1771 mg/kg/day)

Conclusion : Not a teratogen.

Reliability : (2) valid with restrictions

2 - Reliable study with restrictions. No circumstances occurred that would have affected the quality or integrity of the data. Comparable to a guideline study and test procedures were in accordance with generally accepted

scientific standards and described in sufficient detail.

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5.8.3 TOXICITY TO REPRODUCTION, OTHER STUDIES

5.9 SPECIFIC INVESTIGATIONS

5.10 EXPOSURE EXPERIENCE

Remark: Excessive exposure by inhalation may result in headache,

dizziness, drowsiness and narcosis. No adverse systemic effects due to exposure to 2-butanol have been reported in

man.

Source : EXXON CHEMICAL, Limited Fareham, Hampshire

EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)

22.10.1992

5.11 ADDITIONAL REMARKS

Type: other: Respiratory Irritation

Remark : Respiratory rates of CF-1 mice (mean weight 25 g.) were determined prior

to exposure. SBA was evaporated, diluted with room air and fed into a 3.3 liter chamber at an airflow rate of 18.3 to 26.9 l/min. Each animal was placed in a body plethysmograph attached to the exposure chamber so that the head of the animal protruded into the chamber. The respiratory rate and the relative tidal volume were obtained by attaching a pressure transducer to each plethysmograph. The results were expressed as percentage of pre-exposure values, obtained from a 10-minute control period. The exposure was 30 minutes, followed by a 20-minute recovery period during which the respiratory pattern and respiratory rate were monitored continuously. Respiration rates and tidal volumes were compared between normal and cannulated mice to determine the on-set and extent of respiratory irritation. In the cannulated mice, the onset of the decrease in respiration was slower than that of the normal mice indicating

that sBA was acting as a respiratory irritant.

Species/Strain: Mouse / CF-1

Sex: Male

Number/Sex/Dose: 10 Vehicle: None

Route of Administration: Inhalation

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Doses: 2800, 5600, 10100 or 15300 ppm normal mice;

10500 or 14000 ppm cannulated mice

Time of Exposure: 30 minute

Post Dose Observation Period: 20 minute

Method: Alarie Test GLP: Unknown Year: 1994

Result: 640 ppm RD0 Threshold Concentration at 2 minutes;

11800 ppm RD50 at 10 minutes

Test substance : sec-Butanol (sBA)

Conclusion : sBA caused a concentration-dependent decrease in respiratory tidal

volume in normal and cannulated mice and acts as a respiratory irritant.

Reliability : (1) valid without restriction

1 - Reliable without Restrictions. No circumstances occurred that would

have affected the quality or integrity of the data.

11.01.2002 (48)

6. Analyt. Meth. for Detection and Identification	ld 78-92-2 Date
6.1 ANALYTICAL METHODS	
6.2 DETECTION AND IDENTIFICATION	
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7. Eff	. Against Target Org. and Intended Uses	ld	78-92-2
		Date	01.06.2006
7.1	FUNCTION		
7.2	EFFECTS ON ORGANISMS TO BE CONTROLLED		
7.3	ORGANISMS TO BE PROTECTED		
7.4	USER		
7.5	RESISTANCE		
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Id 78-92-2 8. Meas. Nec. to Prot. Man, Animals, Environment Date 01.06.2006 8.1 METHODS HANDLING AND STORING 8.2 FIRE GUIDANCE **EMERGENCY MEASURES** 8.3 POSSIB. OF RENDERING SUBST. HARMLESS 8.4 **WASTE MANAGEMENT SIDE-EFFECTS DETECTION** 8.6 SUBSTANCE REGISTERED AS DANGEROUS FOR GROUND WATER 8.7 8.8 REACTIVITY TOWARDS CONTAINER MATERIAL 70 / 76

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10.1 END POINT SUMMARY	
40.2 HAZADD CHMMADV	
10.2 HAZARD SUMMARY	
10.3 RISK ASSESSMENT	
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    IUCLID-Export V4.00
С
CS
   ISO-Latin 1
С
NL
    GBR
С
B005 SUBST MASTER TAB
F001 6863-58-7
F002 Y26-001
EOB
С
B006 SUBST IDENT TAB
F001 6863-58-7
F002 Y28-001
F003 Y27-001
F004 6863-58-7
F005 1
EOR
F001 6863-58-7
F002 Y28-002
F003 Y27-006
F004 2,2'-oxybisbutane
F005 2
EOR
F001 6863-58-7
F002 Y28-001
F003 Y27-002
F004 229-961-6
F005 3
EOR
F001 6863-58-7
F002 Y28-002
F003 Y27-017
F004 sec-Butyl Ether
F005 4
EOR
F001 6863-58-7
F002 Y28-003
F003 Y27-003
F004 C8H18O
F005 102
EOB
B003 DS ADMIN TAB
F002 520
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F001 6863-58-7
F009 N
F005 12032693
F006 16-10-2006
F007 12032693
F008 16-10-2006
F003 14-11-2006
F101 ExxonMobil Chemical Company - HPV
F102 A35-02
EOB
B004 COMPANY_TAB
F001 12032693
F003 ExxonMobil Biomedical Sciences Inc.
F004 1545 Route 22 East
F005 Annadale, New Jersey
F006 08801-3059
F008 A31-024
EOB
С
С
     ****** N E W D A T A S E T ******
С
D
    520
B052 DS_COMPONENT_JOIN_TAB
F001 520
F002 0
F003 1.1.0
F004 1
F005 1
F006 07-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 1.1.1
F004 1
F005 1
F006 08-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 1.12
F004 1
F005 1
F006 08-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 1.12
F004 2
F005 2
F006 08-11-2006
F007 08-11-2006
EOR
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F001 520
F002 0
F003 1.12
F004 3
F005 3
F006 08-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 1.2
F004 1
F005 1
F006 07-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 1.2
F004 2
F005 2
F006 07-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 1.2
F004 3
F005 3
F006 07-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.1
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.1
F004 2
F005 2
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.2
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
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F002 0
F003 2.3
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.4
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.4
F004 2
F005 2
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.5
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.5
F004 2
F005 2
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.6.1
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 2.6.1
F004 2
F005 2
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
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F003 3.1.1
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 3.1.1
F004 2
F005 2
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 3.1.2
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 3.3.1
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 3.3.1
F004 2
F005 2
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 3.5
F004 1
F005 1
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 3.5
F004 2
F005 2
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 3.7
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F004 1
F005 1
F006 14-11-2006
F007 06-11-2006
EOR
F001 520
F002 0
F003 3.7
F004 2
F005 2
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 4.1
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 4.1
F004 2
F005 2
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 4.1
F004 3
F005 3
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 4.2
F004 1
F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 4.2
F004 2
F005 2
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 4.3
F004 1
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F005 1
F006 14-11-2006
F007 07-11-2006
EOR
F001 520
F002 0
F003 4.3
F004 2
F005 2
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 4.5.1
F004 1
F005 1
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 4.5.1
F004 2
F005 2
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 4.5.2
F004 1
F005 1
F006 14-11-2006
F007 08-11-2006
EOR
F001 520
F002 0
F003 4.5.2
F004 2
F005 2
F006 14-11-2006
F007 08-11-2006
EOB
C
B053 DS_REC_MARK_TAB
F001 520
F002 2.1
F003 1
F004 A37-009
EOR
F001 520
F002 2.2
F003 1
F004 A37-009
EOR
F001 520
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F002 2.3
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F003 1

F004 A37-009

EOR

F001 520

F002 2.4

F003 1

F004 A37-009

EOR

F001 520

F002 2.5

F003 1

F004 A37-009

EOR

F001 520

F002 2.6.1

F003 1

F004 A37-009

EOR

F001 520

F002 3.1.1

F003 1

F004 A37-009

EOR

F001 520

F002 3.1.1

F003 2

F004 A37-009

EOR

F001 520

F002 3.1.2

F003 1

F004 A37-009

EOR

F001 520

F002 3.3.1

F003 1

F004 A37-009

EOR

F001 520

F002 3.3.1

F003 2

F004 A37-009

EOR

F001 520

F002 3.5

F003 1

F004 A37-009

EOR

F001 520

F002 3.7

F003 1

F004 A37-009

EOR

F001 520

F002 4.1

F003 1

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F004 A37-009
EOR
F001 520
F002 4.2
F003 1
F004 A37-009
EOR
F001 520
F002 4.3
F003 1
F004 A37-009
EOR
F001 520
F002 4.5.1
F003 1
F004 A37-009
EOR
F001 520
F002 4.5.2
F003 1
F004 A37-009
EOB
B051 DS_COMPONENT_TAB
F001 520
F002 0
F003 6863-58-7
F012 N
F010 08-11-2006
F004 12032693
F005 16-10-2006
F006 12032693
F007 16-10-2006
F008 ExxonMobil Chemical Company - HPV
F009 A35-02
EOB
B007 GI SUBSTANCE TAB
F001 520
F002 1
F003 07-11-2006
F004 RADAVI
F007 O(C(CC)C)C(CC)C
F008 C8H18O1
F009 130.23
F011 Butane, 2,2'-oxybis-
EOB
B101 GI_GENERAL_INFORM_TAB
F001 520
F002 1
F003 08-11-2006
F004 RADAVI
F010 A04-04
F014 C02-001
EOB
C
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B102 GI_SYNONYM_TAB
F001 520
F002 1
F003 07-11-2006
F004 RADAVI
F007 sec-Butyl Ether
EOR
F001 520
F002 2
F003 07-11-2006
F004 RADAVI
F007 bis (2-butyl) ether
EOR
F001 520
F002 3
F003 07-11-2006
F004 RADAVI
F007 di-sec-Butyl Ether
EOB
B125 GI_LIT_SEARCH_TAB
F001 520
F002 1
F003 08-11-2006
F004 RADAVI
F006 1
F007 A44-003
F009 16-10-2006
EOR
F001 520
F002 2
F003 08-11-2006
F004 RADAVI
F006 2
F007 A44-003
F009 25-05-2005
EOR
F001 520
F002 3
F003 08-11-2006
F004 RADAVI
F006 3
F007 A44-003
F009 20-05-2005
EOB
B201 PC_MELTING_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F015 A36-003
F016 1
F007 A02-03
F008 -100
F012 P01-03: measured
F014 A03-02
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F020 A01-03: CAS No. 6863-58-7; sec-butyl ether
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F015 A36-003
F016 2
F007 A02-03
F008 -73
F012 P01-03: calculated
F013 2003
F014 A03-01
F020 A01-03: CAS No. 6863-58-7; sec-butyl ether
EOB
B202 PC_BOILING_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F016 A36-003
F017 1
F007 A02-03
F008 116
F010 1013
F011 P02-01
F013 P03-03: calculated
F014 2003
F015 A03-01
F018 A01-03: CAS No. 6863-58-7; sec-butyl ether
EOB
B203 PC_DENSITY_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F016 A36-003
F017 1
F008 A02-03
F009 .759
F011 P18-01
F012 25
F013 P04-03: measured
F014 1935
F015 A03-02
F018 A01-03: CAS No. 6863-58-7; sec-butyl ether
EOB
B204 PC_VAPOUR_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F015 A36-003
F016 1
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F007 A02-03
F008 21.7
F010 P02-01
F011 25
F012 P06-04
F013 1994
F014 A03-02
F018 A01-03: CAS No. 6863-58-7; sec-butyl ether
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F015 A36-003
F016 2
F007 A02-03
F008 29.7
F010 P02-01
F011 25
F012 P06-03
F013 2003
F014 A03-01
F018 A01-03: CAS No. 6863-58-7; sec-butyl ether
EOB
B205 PC_PARTITION_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F014 A36-003
F015 1
F007 A02-03
F008 3.35
F010 25
F011 P07-05
F012 1992
F013 A03-02
F016 A01-03: CAS No. 6863-58-7; sec-butyl ether
F020 C15-001
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F014 A36-003
F015 2
F007 A02-03
F008 2.87
F010 25
F011 P07-04
F012 2003
F013 A03-01
F016 A01-03: CAS No. 6863-58-7; sec-butyl ether
F020 C15-001
EOB
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B206 PC_WATER_SOL_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F023 A36-003
F024 1
F007 A02-03
F008 P08-02
F009 330
F011 25
F020 P09-03: measured
F021 1992
F022 A03-02
F025 A01-03: CAS No. 6863-58-7; sec-butyl ether
F030 C14-001
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F023 A36-003
F024 2
F007 A02-03
F008 P08-02
F009 327
F011 25
F020 P09-03: calculated
F021 2003
F022 A03-01
F025 A01-03: CAS No. 6863-58-7; sec-butyl ether
F030 C14-001
EOB
С
B301 EN_PHOTODEGRADATION_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F045 A36-003
F046 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 F01-01
F009 F02-05: Calculated values using AOPWIN version 1.91, a subroutine of the
     computer program EPIWIN version 3.12
F010 2003
F011 F03-01
F034 F06-03
F035 1500000
F036 F07-02
F044 A02-03
F037 .00000000000320784
F043 A03-01
EOR
F001 520
F002 2
F003 14-11-2006
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F004 CLGETTS1
F046 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 F01-04
F009 F02-05: Technical discussion
F010 2006
EOB
B302 EN_STABILITY_IN_WATER_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F041 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 F08-01
F009 F09-03: technical discussion
F010 2006
EOB
B305 EN_TRANSPORT_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F011 A36-003
F012 1
F007 F20-07
F008 F22-01: air - sediment(s) - soil - water
F009 F21-01: Calculation according Mackay, Level III
F010 2006
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F011 A36-003
F012 2
F007 F20-05
F008 F22-01: air - biota - sediment(s) - soil - water
F009 F21-01: Calculation according Mackay, Level I
F010 2006
EOB
B308 EN BIODEGRADATION TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F047 A36-003
F048 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 F25-01
F009 F26-25: calculated using BIOWIN version 4.02
F020 F30-02: not readily biodegradable
F046 A03-01
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EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F047 A36-003
F048 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 F25-01
F009 F26-25: Japanese Guideline by MITI (1974). Comparable to OECD TG 301C,
    Modified MITI Test 1
F010 1992
F011 F27-0137
F012 100
F013 F28-02
F014 F29-03
F015 A02-03
F016 4
F017 3
F018 28
F019 F05-01
F020 F30-02: not readily biodegradable
F046 A03-02
EOB
B310 EN_BIOACCUMULATION_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F021 A36-003
F022 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E02-0033
F009 F34-06: guideline corresponds to OECD 305C, Degree of Bioconcentration in
     Fish (1981)
F011 42
F012 F10-01
F016 A02-03
F017 47
F018 83
F020 A03-02
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F021 A36-003
F022 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F009 F34-06: calculated
F010 2006
F016 A02-03
F017 32.1
F020 A03-01
EOB
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B401 EC_FISHTOX_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F033 A36-003
F034 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E01-03: calculated
F009 E02-0161: freshwater fish
F010 E03-05: ECOSAR Computer Model
F011 2006
F012 96
F013 E04-02
F014 E05-02
F021 A02-03
F022 14.7
F045 E35-01
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F033 A36-003
F034 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E01-03: calculated
F009 E02-0161: freshwater fish
F010 E03-05: ECOSAR Computer Model
F011 2006
F012 96
F013 E04-02
F014 E05-02
F021 A02-03
F022 10.8
F032 A03-01
F045 E35-01
EOR
F001 520
F002 3
F003 14-11-2006
F004 CLGETTS1
F033 A36-003
F034 3
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E01-04
F009 E02-0106
F010 E03-05: Japanese Industrial Standard (JIS K 0102-1986-71): Testing
     Methods for Industrial Wastewater.
F011 1992
F012 48
F013 E04-02
F014 E05-02
F021 A02-03
F022 30.7
F032 A03-02
F045 E35-02
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F050 C47-001
EOB
B402 EC_DAPHNIATOX_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F032 A36-003
F033 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E06-0013
F009 E07-04: ECOSAR Computer Model
F010 2006
F011 48
F012 E04-02
F013 E05-02
F020 A02-03
F021 16.7
F031 A03-01
F045 E35-01
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F032 A36-003
F033 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E06-0013
F009 E07-04: ECOSAR Computer Model
F010 2006
F011 48
F012 E04-02
F013 E05-02
F020 A02-03
F021 12.5
F031 A03-01
F045 E35-01
EOB
B403 EC_ALGAETOX_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F036 A36-003
F037 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E08-0063: green alga
F009 E09-04: ECOSAR Computer Model
F010 2006
F012 96
F013 E04-02
F014 E05-02
F027 A02-03
F028 11
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F035 A03-01
F050 E35-01
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F036 A36-003
F037 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E08-0063: green alga
F009 E09-04: ECOSAR Computer Model
F010 2006
F012 96
F013 E04-02
F014 E05-02
F027 A02-03
F028 8.3
F035 A03-01
F050 E35-01
EOB
B405 EC_CHRONFISHTOX_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F030 A36-003
F031 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E02-0161: Freshwater Fish (calculated toxicity values are not species
     specific)
F009 E13-02: ECOSAR Computer Model
F010 2006
F011 E14-02: LC50
F012 30
F013 E15-01
F014 E05-02
F024 ChV
F025 A02-03
F026 2.2
F029 A03-01
F035 E35-01
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F030 A36-003
F031 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E02-0161: Freshwater Fish (calculated toxicity values are not species
     specific)
F009 E13-02: ECOSAR Computer Model
F010 2006
F011 E14-02: LC50
F012 30
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F013 E15-01
F014 E05-02
F024 ChV
F025 A02-03
F026 1.6
F029 A03-01
F035 E35-01
EOB
B406 EC_CHRONDAPHNIATOX_TAB
F001 520
F002 1
F003 14-11-2006
F004 CLGETTS1
F030 A36-003
F031 1
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E06-0013
F009 E16-02: ECOSAR Computer model
F010 2006
F011 E17-01
F012 16
F013 E18-01
F014 E05-02
F015 A02-03
F016 1.3
F029 A03-01
F032 E35-01
EOR
F001 520
F002 2
F003 14-11-2006
F004 CLGETTS1
F030 A36-003
F031 2
F007 A01-03: CAS No. 6863-58-7; sec-butyl ether
F008 E06-0013
F009 E16-02: ECOSAR Computer model
F010 2006
F011 E17-01
F012 16
F013 E18-01
F014 E05-02
F015 A02-03
F016 1
F029 A03-01
F032 E35-01
EOB
B601 TEXT_TAB
F002 520
F010 1.12
F004 1
F005 RM
F006 Search covered all Physical Chemical Properties, Environmental Fate,
     Aquatic and Mammalian Toxicity endpoints related to the CAS number.
F007 Search covered all Physical Chemical Properties, Environmental Fate,
```

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Aquatic and Mammalian Toxicity endpoints related to the CAS number.
F020 260662
EOR
F002 520
F010 1.12
F004 2
F005 RM
F006 Search covered all Physical Chemical Properties, Environmental Fate,
     Aquatic and Mammalian Toxicity endpoints related to the CAS number.
F007 Search covered all Physical Chemical Properties, Environmental Fate,
     Aquatic and Mammalian Toxicity endpoints related to the CAS number.
F020 260663
EOR
F002 520
F010 1.12
F004 3
F005 RM
F006 Search covered all Physical Chemical Properties, Environmental Fate,
     Aquatic and Mammalian Toxicity endpoints related to the CAS number.
F007 Search covered all Physical Chemical Properties, Environmental Fate,
     Aquatic and Mammalian Toxicity endpoints related to the CAS number.
F020 260664
EOR
F002 520
F010 2.1
F004 1
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260558
EOR
F002 520
F010 2.1
F004 1
F005 RL
F006 Although the original data were not retrieved and reviewed for quality,
     they were developed following acceptable test methods and therefore
     considered reliable. Value was provided by the experimental database of
     the EPIWIN program.
F007 Although the original data were not retrieved and reviewed for quality,
     they were developed following acceptable test methods and therefore
     considered reliable. Value was provided by the experimental database of
     the EPIWIN program.
F020 260557
EOR
F002 520
F010 2.1
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260546
EOR
F002 520
F010 2.1
```

```
F004 2
F005 ME
F006 Calculated values using MPBPWIN version 1.41, a subroutine of the
     computer program EPIWIN version 3.12
F007 Calculated values using MPBPWIN version 1.41, a subroutine of the
     computer program EPIWIN version 3.12
F020 260573
EOR
F002 520
F010 2.1
F004 2
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260572
EOR
F002 520
F010 2.1
F004 2
F005 RL
F006 The result is a calculated value based on the chemical structure and
     represents a potential melting point for the substance with the CAS
    number listed under test substance.
F007 The result is a calculated value based on the chemical structure and
     represents a potential melting point for the substance with the CAS
     number listed under test substance.
F020 260575
EOR
F002 520
F010 2.1
F004 2
F005 TC
F006 Melting Point estimations performed by MPBPWIN are based on the average
     result of the calculation methods of K. Joback and Gold and Ogle.
* *
* *
     Joback's Method is described in Joback, K.G. 1982. A Unified Approach to
     Physical Property Estimation
F007 Melting Point estimations performed by MPBPWIN are based on the average
    result of the calculation methods of K. Joback and Gold and Ogle.
* *
     Joback's Method is described in Joback, K.G. 1982. A Unified Approach to
     Physical Property Estimation Using Multivariate Statistical Techniques.
     In The Properties of Gases and Liquids. Fourth Edition. 1987. R.C. Reid,
    J.M. Prausnitz and B.E. Poling, Eds.
* *
* *
    The Gold and Ogle Method simply uses the formula
     Tm = 0.5839Tb, where Tm is the melting point in Kelvin and Tb is the
     boiling point in Kelvin.
F020 260574
EOR
F002 520
F010 2.1
F004 2
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
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F007 CAS No. 6863-58-7; sec-butyl ether
F020 260571
EOR
F002 520
F010 2.2
F004 1
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260559
EOR
F002 520
F010 2.2
F004 1
F005 RL
F006 The result is a calculated value based on the chemical structure and
     represents a potential boiling point for the substance with the CAS
    number listed under test substance.
F007 The result is a calculated value based on the chemical structure and
     represents a potential boiling point for the substance with the CAS
     number listed under test substance.
F020 260561
EOR
F002 520
F010 2.2
F004 1
F005 TC
F006 Boiling point calculated by MPBPWIN subroutine, which is based on the
     method of S. Stein and R. Brown in "Estimation of Normal Boiling Points
     from Group Contributions". 1994. J. Chem. Inf. Comput. Sci. 34: 581-587.
F007 Boiling point calculated by MPBPWIN subroutine, which is based on the
     method of S. Stein and R. Brown in "Estimation of Normal Boiling Points
     from Group Contributions". 1994. J. Chem. Inf. Comput. Sci. 34: 581-587.
F020 260560
EOR
F002 520
F010 2.2
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260547
EOR
F002 520
F010 2.3
F004 1
F005 RE
F006 Drake, Nathan L. (1935). Journal of the American Chemical Society. Vol
     57, pp. 2623-2625 CAPLUS.
F007 Drake, Nathan L. (1935). Journal of the American Chemical Society. Vol
     57, pp. 2623-2625 CAPLUS.
F020 260562
EOR
F002 520
F010 2.3
```

```
F004 1
F005 RL
F006 Although the original data were not retrieved and reviewed for quality,
     they were reported in the peer-reviewed literature and therefore
     considered reliable.
F007 Although the original data were not retrieved and reviewed for quality,
     they were reported in the peer-reviewed literature and therefore
     considered reliable.
F020 260563
EOR
F002 520
F010 2.3
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260548
EOR
F002 520
F010 2.4
F004 1
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260565
EOR
F002 520
F010 2.4
F004 1
F005 RL
F006 Although the original data were not retrieved and reviewed for quality,
     they were developed following acceptable test methods and therefore
     considered reliable. Value was provided by the experimental database of
     the EPIWIN program.
* *
* *
     Referen
F007 Although the original data were not retrieved and reviewed for quality,
     they were developed following acceptable test methods and therefore
     considered reliable. Value was provided by the experimental database of
     the EPIWIN program.
* *
* *
     Reference given in EPI Suite: Yaws, CL (1994B).
F020 260564
EOR
F002 520
F010 2.4
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260549
EOR
F002 520
F010 2.4
F004 2
```

```
F005 ME
F006 Vapor pressure calculation by MPBPWIN ver. 1.40 using calculation method
     of Grain.
F007 Vapor pressure calculation by MPBPWIN ver. 1.40 using calculation method
     of Grain.
F020 260567
EOR
F002 520
F010 2.4
F004 2
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260570
EOR
F002 520
F010 2.4
F004 2
F005 RL
F006 The result is a calculated value based on the chemical structure and
     represents a potential vapor pressure for the substance with the CAS
     number listed under test substance.
F007 The result is a calculated value based on the chemical structure and
     represents a potential vapor pressure for the substance with the CAS
     number listed under test substance.
F020 260569
EOR
F002 520
F010 2.4
F004 2
F005 RM
F006 EPIWIN is used and advocated by the US EPA for chemical property
     estimation.
F007 EPIWIN is used and advocated by the US EPA for chemical property
     estimation.
F020 260568
EOR
F002 520
F010 2.4
F004 2
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260566
EOR
F002 520
F010 2.5
F004 1
F005 RE
F006 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
     CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
     Ministry of Internatio
F007 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
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and bioaccumulation data of existing chemicals based on the CSCL Japan.

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CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
    Ministry of International Trade and Industry, Japan. Japan Chemical
     Industry Ecology-Toxicology and Information Center.
F020 260576
EOR
F002 520
F010 2.5
F004 1
F005 RL
F006 Although the original data were not retrieved and reviewed for quality,
     they were developed at a well respected testing facility and therefore
     considered reliable.
F007 Although the original data were not retrieved and reviewed for quality,
     they were developed at a well respected testing facility and therefore
     considered reliable.
F020 260583
EOR
F002 520
F010 2.5
F004 1
F005 TC
F006 Specific methods were not given in the document reporting the measured
     value. Measurements were performed at the Chemicals Inspection and
    Testing Institute of Japan.
F007 Specific methods were not given in the document reporting the measured
     value. Measurements were performed at the Chemicals Inspection and
     Testing Institute of Japan.
F020 260582
EOR
F002 520
F010 2.5
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260550
EOR
F002 520
F010 2.5
F004 2
F005 ME
F006 Calculated values using KOWWIN version 1.67, a subroutine of the computer
     program EPIWIN version 3.12
F007 Calculated values using KOWWIN version 1.67, a subroutine of the computer
     program EPIWIN version 3.12
F020 260578
EOR
F002 520
F010 2.5
F004 2
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260581
EOR
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F002 520
F010 2.5
F004 2
F005 RL
F006 The result is a calculated value based on the chemical structure and
     represents a potential partition coefficient for the substance with the
     CAS number listed under test substance.
F007 The result is a calculated value based on the chemical structure and
     represents a potential partition coefficient for the substance with the
     CAS number listed under test substance.
F020 260580
EOR
F002 520
F010 2.5
F004 2
F005 TC
F006 Octanol / Water Partition Coefficient estimations performed by KOWWIN are
     based on an atom/fragment contribution method of W. Meylan and P. Howard
     in "Atom/fragment contribution method for estimating octanol-water
    partition coefficients". 1
F007 Octanol / Water Partition Coefficient estimations performed by KOWWIN are
    based on an atom/fragment contribution method of W. Meylan and P. Howard
     in "Atom/fragment contribution method for estimating octanol-water
     partition coefficients". 1995. J. Pharm. Sci. 84:83-92.
F020 260579
EOR
F002 520
F010 2.5
F004 2
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260577
EOR
F002 520
F010 2.6.1
F004 1
F005 RE
F006 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
     CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
    Ministry of Internatio
F007 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
     CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
     Ministry of International Trade and Industry, Japan. Japan Chemical
     Industry Ecology-Toxicology and Information Center.
F020 260584
EOR
F002 520
F010 2.6.1
F004 1
F005 RL
F006 Although the original data were not retrieved and reviewed for quality,
     they were developed at a well respected testing facility and therefore
     considered reliable.
F007 Although the original data were not retrieved and reviewed for quality,
```

```
they were developed at a well respected testing facility and therefore
     considered reliable.
F020 260585
EOR
F002 520
F010 2.6.1
F004 1
F005 TC
F006 Specific methods were not given in the document reporting the measured
     value. Measurements were performed at the Chemicals Inspection and
    Testing Institute of Japan.
F007 Specific methods were not given in the document reporting the measured
     value. Measurements were performed at the Chemicals Inspection and
     Testing Institute of Japan.
F020 260586
EOR
F002 520
F010 2.6.1
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260551
EOR
F002 520
F010 2.6.1
F004 2
F005 ME
F006 Calculated values using WSKOWWIN version 1.41, a subroutine of the
     computer program EPIWIN version 3.12
F007 Calculated values using WSKOWWIN version 1.41, a subroutine of the
     computer program EPIWIN version 3.12
F020 260588
EOR
F002 520
F010 2.6.1
F004 2
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260591
EOR
F002 520
F010 2.6.1
F004 2
F005 RL
F006 The result is a calculated value based on the chemical structure and
     represents a potential water solubility for the substance with the CAS
     number listed under test substance.
F007 The result is a calculated value based on the chemical structure and
    represents a potential water solubility for the substance with the CAS
     number listed under test substance.
F020 260590
EOR
```

F002 520

```
F010 2.6.1
F004 2
F005 TC
F006 Water Solubility estimations performed by WSKOWWIN are based on a Kow
     correlation method described by W. Meylan, P. Howard and R. Boethling in
     "Improved method for estimating water solubility from octanol/water
    partition coefficient". Envir
F007 Water Solubility estimations performed by WSKOWWIN are based on a Kow
     correlation method described by W. Meylan, P. Howard and R. Boethling in
     "Improved method for estimating water solubility from octanol/water
     partition coefficient". Environ. Toxicol. Chem. 15:100-106. 1995.
F020 260589
EOR
F002 520
F010 2.6.1
F004 2
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260587
EOR
F002 520
F010 3.1.1
F004 1
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260593
EOR
F002 520
F010 3.1.1
F004 1
F005 RL
F006 The results include calculated data based on chemical structure as
     modeled by AOPWIN. The data represent a potential atmospheric half-life
    range for the test substance.
F007 The results include calculated data based on chemical structure as
    modeled by AOPWIN. The data represent a potential atmospheric half-life
     range for the test substance.
F020 260596
E \cap R
F002 520
F010 3.1.1
F004 1
F005 RS
F006 Atmospheric Oxidation Potential
     In the environment, organic chemicals emitted into the troposphere are
     degraded by several important transformation processes. The dominant
     transformation process for most compounds is the daylight reaction
F007 Atmospheric Oxidation Potential
* *
     In the environment, organic chemicals emitted into the troposphere are
    degraded by several important transformation processes. The dominant
```

transformation process for most compounds is the daylight reaction with

```
hydroxyl (OH-) radicals (Atkinson, 1988, 1989). The rate at which an
     organic compound reacts with OH- radicals is a direct measure of its
     atmospheric persistence (Meylan and Howard, 1993).
* *
     AOPWIN estimates the rate constant for the atmospheric, gas-phase
     reaction between photochemically produced hydroxyl radicals and organic
     chemicals. The rate constants estimated by the program are then used to
     calculate atmospheric half-lives for organic compounds based upon average
     atmospheric concentrations of hydroxyl radicals.
* *
     Since the reactions only take place in the presence of sunlight, the
     atmospheric half-lives are normalized for a 12-hour day.
* *
     Calculated*
                        OH- Rate Constant
* *
     half-life (days)
                              (cm3/molecule-sec)
* *
* *
     0.333
                        32.0784 E-12
* *
* *
     References:
* *
* *
     Atkinson, R. 1988. Estimation of gas-phase hydroxyl radical rate
     constants for organic chemicals. Environ. Toxicol. Chem. 7:435-442.
* *
* *
     Atkinson, R. 1989. Kinetics and mechanisms of the gas-phase reactions of
     the hydroxyl radical with organic compounds. J. Phys. Chem. Ref. Data
     Monograph No. 1, Amer. Inst. Physics & Amer. Chem. Soc., NY.
* *
* *
     Meylan, W.M. and P.H. Howard. 1993. Computer estimation of the
     atmospheric gas-phase reaction rate of organic compounds with hydroxyl
     radicals and ozone. Chemosphere 12:2293-2299.
F020 260594
EOR
F002 520
F010 3.1.1
F004 1
F005 TC
F006
* *
     Indirect photodegradation, or atmospheric oxidation potential, is based
     on the structure-activity relationship methods developed by R. Atkinson.
* *
* *
     Temperature: 25°C
* *
     Sensitizer: OH radical
* *
     Concentration of Sensitizer:
                                    1.5 E6 OH radicals/cm3
F007
* *
     Indirect photodegradation, or atmospheric oxidation potential, is based
     on the structure-activity relationship methods developed by R. Atkinson.
* *
* *
     Temperature: 25°C
* *
     Sensitizer: OH radical
* *
     Concentration of Sensitizer: 1.5 E6 OH radicals/cm3
F020 260595
F002 520
F010 3.1.1
```

```
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260592
EOR
F002 520
F010 3.1.1
F004 2
F005 RE
F006 EMBSI (2006) Photodegradation (Direct): CAS No. 6863-58-7; sec-Butyl
     Ether.
F007 EMBSI (2006) Photodegradation (Direct): CAS No. 6863-58-7; sec-Butyl
     Ether.
F020 260599
EOR
F002 520
F010 3.1.1
F004 2
F005 RS
F006 Photolysis as a Function of Molecular Structure
* *
     The direct photolysis of an organic molecule occurs when it absorbs
     sufficient light energy to result in a structural transformation (Harris,
     1982). The reaction process is initiated when lig
F007 Photolysis as a Function of Molecular Structure
* *
* *
     The direct photolysis of an organic molecule occurs when it absorbs
     sufficient light energy to result in a structural transformation (Harris,
     1982). The reaction process is initiated when light energy in a specific
     wavelength range elevates a molecule to an electronically excited state.
    However, the excited state is competitive with various deactivation
    processes that can result in the return of the molecule to a non excited
     state.
* *
     The absorption of light in the ultra violet (UV)-visible range, 110-750
     nm, can result in the electronic excitation of an organic molecule. Light
     in this range contains energy of the same order of magnitude as covalent
    bond dissociation energies (Harris, 1982). Higher wavelengths (e.g.
     infrared) result only in vibrational and rotational transitions, which do
    not tend to produce structural changes to a molecule.
    The stratospheric ozone layer prevents UV light of less than 290 nm from
     reaching the earth's surface. Therefore, only light at wavelengths
     between 290 and 750 nm can result in photochemical transformations in the
     environment (Harris, 1982). Although the absorption of UV light in the
     290-750 nm range is necessary, it is not always sufficient for a chemical
     to undergo photochemical degradation. Energy may be re-emitted from an
     excited molecule by mechanisms other than chemical transformation,
     resulting in no change to the parent molecule.
* *
    A conservative approach to estimating a photochemical degradation rate is
     to assume that degradation will occur in proportion to the amount of
     light wavelengths >290 nm absorbed by the molecule (Zepp and Cline,
     1977).
* *
```

Saturated hydrocarbons and R-OH groups do not absorb light above 290 nm

* *

```
(Harris J, 1982a). Therefore, these moieties are stable in regard to
     direct photolytic processes. Ethers are also stable as this group absorbs
     UV light in the far UV region, below 220 nm (Mill T, 2000), therefore,
     direct photolysis will not be an important transformation process for the
     degradation of sec-butyl ether in the environment.
* *
* *
     References:
* *
* *
    Harris, J. C. 1982. "Rate of Aqueous Photolysis," Chapter 8 in: W. J.
    Lyman, W. F. Reehl, and D. H. Rosenblatt, eds., Handbook of Chemical
     Property Estimation Methods, McGraw-Hill Book Company, New York, USA.
* *
     Zepp, R. G. and D. M. Cline. 1977. Rates of Direct Photolysis in the
     Aqueous Environment, Environ. Sci. Technol., 11:359-366.
* *
* *
     Boethling, R.S., Mackay, D. 2000. Handbook of Property Estimation
     Methods for Chemicals, CRC Press, Boca Raton, FL, USA.
F020 260598
EOR
F002 520
F010 3.1.1
F004 2
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260597
EOR
F002 520
F010 3.1.2
F004 1
F005 RE
F006 EMBSI (2006) Hydrolysis: CAS No. 6863-58-7; sec-Butyl Ether.
F007 EMBSI (2006) Hydrolysis: CAS No. 6863-58-7; sec-Butyl Ether.
F020 260602
EOR
F002 520
F010 3.1.2
F004 1
F005 RS
F006 Hydrolysis as a Function of Molecular Structure
* *
     Hydrolysis of an organic molecule occurs when a molecule (R-X) reacts
     with water (H2O) to form a new carbon-oxygen bond after the carbon-X bond
     is cleaved (Gould, 1959; Harris, 1982). Mechani
F007 Hydrolysis as a Function of Molecular Structure
* *
     Hydrolysis of an organic molecule occurs when a molecule (R-X) reacts
     with water (H2O) to form a new carbon-oxygen bond after the carbon-X bond
     is cleaved (Gould, 1959; Harris, 1982). Mechanistically, this reaction is
     referred to as a nucleophilic substitution reaction, where X is the
     leaving group being replaced by the incoming nucleophilic oxygen from the
    water molecule.
    Chemicals that are susceptible to hydrolysis contain functional groups
     that can be displaced by a nucleophilic substitution reaction. Substances
```

that have a potential to hydrolyze include alkyl halides, amides, carbamates, carboxylic acid esters and lactones, epoxides, phosphate

```
esters, and sulfonic acid esters (Neely, 1985). The lack of a suitable
     leaving group renders a compound resistant to hydrolysis.
* *
* *
     Sec-butyl ether is expected to be resistant to hydrolysis because it
     lacks a functional group that is hydrolytically reactive (Harris, 1982b).
      Therefore, hydrolysis will not contribute to its removal from the
     environment.
* *
* *
     References:
* *
* *
     Gould, E.S. (1959), Mechanism and Structure in Organic Chemistry, Holt,
     Reinhart and Winston, New York, NY, USA.
* *
* *
     Harris, J.C. (1982), "Rate of Hydrolysis," Chapter 7 in: W.J. Lyman, W.F.
     Reehl, and D.H. Rosenblatt, eds., Handbook of Chemical Property
     Estimation Methods, McGraw-Hill Book Company, New York, NY, USA.
* *
* *
     Neely, W. B. 1985. Hydrolysis. In: W. B. Neely and G. E. Blau, eds.
     Environmental Exposure from Chemicals. Vol I., pp. 157-173. CRC Press,
     Boca Raton, FL, USA.
F020 260601
EOR
F002 520
F010 3.1.2
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260600
EOR
F002 520
F010 3.3.1
F004 1
F005 ME
F006 The EQC Level III model is a steady state model that is useful for
     determining how the medium of release affects environmental fate. Level
     III fugacity allows non-equilibrium conditions to exist between connected
     media as steady state, and
F007 The EQC Level III model is a steady state model that is useful for
     determining how the medium of release affects environmental fate. Level
     III fugacity allows non-equilibrium conditions to exist between connected
     media as steady state, and illustrate important transport and
     transformation processes.
* *
* *
     Physicochemical input values for the model were calculated using the
     EPIWIN Estimation v 3.12 program. Measured input values were also used
     where available and obtained from the EPIWIN database or other reliable
     sources. Distribution data from the equilibrium model provide basic
     information on the potential partitioning behavior of chemicals between
     selected environmental compartments (i.e., air, water, soil, and
     sediment).
* *
     Input values used:
* *
     Molecular mass = 130.23 g/mol
     Water solubility = 327 mg/L
* *
* *
     Vapour pressure = 2170 Pa
* *
     log Kow = 2.87
```

```
* *
     Melting point = -100 \deg C
* *
* *
     Degradation half-lives:
* *
* *
     Air - 4.0 hrs
* *
     Water - 360 hrs
* *
     Soil - 7200 hrs
* *
     Sediment - 72000 hrs
* *
* *
     This model was run assuming the default emissions.
F020 260603
EOR
F002 520
F010 3.3.1
F004 1
F005 RE
F006 Mackay D, DiGuardo A, Paterson S and Cowan C (2003). EQC Model ver. 2.02,
     available from the Environmental Centre, Trent University, Canada.
F007 Mackay D, DiGuardo A, Paterson S and Cowan C (2003). EQC Model ver. 2.02,
     available from the Environmental Centre, Trent University, Canada.
F020 260606
EOR
F002 520
F010 3.3.1
F004 1
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F020 260605
EOR
F002 520
F010 3.3.1
F004 1
F005 RS
F006 Air - 3.2%
* *
    Water - 44.8%
* *
     Soil - 51.1%
* *
    Sediment - 0.9%
F007 Air - 3.2%
     Water - 44.8%
     Soil - 51.1%
* *
     Sediment - 0.9%
F020 260604
EOR
F002 520
F010 3.3.1
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260552
EOR
F002 520
F010 3.3.1
F004 2
```

```
F005 ME
F006 The EQC Level I is a steady state, equilibrium model that utilizes the
     input of basic chemical properties including molecular weight, vapor
     pressure, and water solubility to calculate distribution within a
     standardized regional environment.
F007 The EQC Level I is a steady state, equilibrium model that utilizes the
     input of basic chemical properties including molecular weight, vapor
     pressure, and water solubility to calculate distribution within a
     standardized regional environment.
* *
     Physicochemical input values for the model were calculated using the
     EPIWIN Estimation v 3.12 program. Measured input values were also used
     where available and obtained from the EPIWIN database or other reliable
     sources. Distribution data from the equilibrium model provide basic
     information on the potential partitioning behavior of chemicals between
     selected environmental compartments (i.e., air, water, soil, and
     sediment).
* *
     Input values used:
* *
     Molecular mass = 130.23 g/mol
* *
     Water solubility = 327 mg/L
* *
     Vapour pressure = 2170 Pa
* *
     log Kow = 2.87
* *
     Melting point = -100 deg C
F020 260608
EOR
F002 520
F010 3.3.1
F004 2
F005 RE
F006 Mackay D, DiGuardo A, Paterson S and Cowan C (2003). EQC Model ver. 2.02,
     available from the Environmental Centre, Trent University, Canada.
F007 Mackay D, DiGuardo A, Paterson S and Cowan C (2003). EQC Model ver. 2.02,
     available from the Environmental Centre, Trent University, Canada.
F020 260607
EOR
F002 520
F010 3.3.1
F004 2
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F020 260610
EOR
F002 520
F010 3.3.1
F004 2
F005 RS
F006 Air - 99.1%
     Water - 0.5%
* *
     Soil - 0.4%
* *
     Sediment - <0.01%
* *
     Suspended Sed - <0.01%
* *
    Biota - <0.01%
F007 Air - 99.1%
```

```
Water - 0.5%
* *
     Soil - 0.4%
* *
     Sediment - <0.01%
* *
     Suspended Sed - <0.01%
* *
    Biota - <0.01%
F020 260609
EOR
F002 520
F010 3.3.1
F004 2
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260553
EOR
F002 520
F010 3.5
F004 1
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260673
EOR
F002 520
F010 3.5
F004 1
F005 RE
F006 Howard, PH, Boethling, RS, Stitler, WM, Meylan, WM, Hueber, AE, Beauman,
     JA and Larosche, ME (1992). Predictive model for aerobic
     biodegradability developed from a file of evaluated biodegradation data.
     Environ Toxicol Chem 11, 593-603.
F007 Howard, PH, Boethling, RS, Stitler, WM, Meylan, WM, Hueber, AE, Beauman,
     JA and Larosche, ME (1992). Predictive model for aerobic
     biodegradability developed from a file of evaluated biodegradation data.
     Environ Toxicol Chem 11, 593-603.
F020 260674
EOR
F002 520
F010 3.5
F004 1
F005 RL
F006 The value was calculated based on the chemical structure as modeled by
     EPI Suite. This robust summary has a reliability rating of 2 because the
     data are modeled.
F007 The value was calculated based on the chemical structure as modeled by
     EPI Suite. This robust summary has a reliability rating of 2 because the
     data are modeled.
F020 260672
EOR
F002 520
F010 3.5
F004 1
F005 RM
F006 Calculation of biodegradation and the timeframe for Primary and Ultimate
     biodegradation using BIOWIN version 4.02, a subroutine of the computer
```

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program EPI Suite version 3.12 as described by Howard, et al, 1994.
* *
     BIOWIN contains six models
F007 Calculation of biodegradation and the timeframe for Primary and Ultimate
     biodegradation using BIOWIN version 4.02, a subroutine of the computer
     program EPI Suite version 3.12 as described by Howard, et al, 1994.
* *
* *
    BIOWIN contains six models (linear regression (BIOWIN 1), non-linear
    regression (BIOWIN 2), ultimate degradation to CO2 and H2O (BIOWIN 3),
     primary degradation (BIOWIN 4), linear regression estimate of the
    probability of passing the OECD 301C / MITI-1 ready biodegradation test
     (BIOWIN 5), and non-linear regression estimate of the probability of
    passing the OECD 301C / MITI-1 ready biodegradation test (BIOWIN 6).
* *
    BIOWIN 1 - "Does Not Biodegrade Fast"
* *
    BIOWIN 2 - "Does Not Biodegrade Fast"
* *
     BIOWIN 3 - "Weeks"
     BIOWIN 4 - "Days-Weeks"
* *
* *
    BIOWIN 5 - "Does Not Biodegrade Fast"
* *
     BIOWIN 6 - "Does Not Biodegrade Fast"
* *
    According to the USEPA, BIOWIN 6 is better predictor of whether a
     chemical will pass or fail the OECD 301C / MITI-1 ready biodegradation
     test than BIOWIN 5.
F020 260671
EOR
F002 520
F010 3.5
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260670
EOR
F002 520
F010 3.5
F004 2
F005 ME
F006 The test was conducted in accordance with "Biodegradation test of
     chemical substances by microorganisms etc". stipulated in the Order
     Prescribing the Items of the Test Relating to the New Chemical Substance
     (1974, Order of the Prime Ministe
F007 The test was conducted in accordance with "Biodegradation test of
     chemical substances by microorganisms etc". stipulated in the Order
     Prescribing the Items of the Test Relating to the New Chemical Substance
     (1974, Order of the Prime Minister, the Minister of Health and Welfare,
     the Minister of International Trade and Industry No 1). This guideline
     coresponds to "301C, Ready Biodegradability: Modified MITI Test 1"
     stipulated in the OECD Guidelines for Testing of Chemicals (1981).
F020 260677
EOR
F002 520
F010 3.5
F004 2
F005 RE
F006 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
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CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
     Ministry of Internatio
F007 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
     CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
     Ministry of International Trade and Industry, Japan. Japan Chemical
     Industry Ecology-Toxicology and Information Center.
F020 260676
EOR
F002 520
F010 3.5
F004 2
F005 RL
F006 Although a standard method was not followed, the testing procedures
     followed generally accepted aerobic biodegradation guideline methods and
     although there was limited information on the specifics of the study,
     there was sufficient informat
F007 Although a standard method was not followed, the testing procedures
     followed generally accepted aerobic biodegradation guideline methods and
     although there was limited information on the specifics of the study,
     there was sufficient information on testing method and conditions in
     general to rate this study a 2.
F020 260679
EOR
F002 520
F010 3.5
F004 2
F005 TC
F006 Sludge concentration = 30 mg/l
F007 Sludge concentration = 30 mg/l
F020 260678
EOR
F002 520
F010 3.5
F004 2
F005 TS
F006 Analogue substance CAS No. 142-96-1; n-butyl ether
F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260675
EOR
F002 520
F010 3.7
F004 1
F005 RE
F006 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
          (ed.). Chemical Products Safety Division, Basic Industries Bureau,
    Ministry of Internatio
F007 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
     CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
    Ministry of International Trade and Industry, Japan. Japan Chemical
     Industry Ecology-Toxicology and Information Center.
F020 260544
EOR
F002 520
F010 3.7
```

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F004 1
F005 RL
F006 Although a standard method was not followed, the testing procedures
     followed generally accepted fish bioconcentration guideline methods and
     although there was limited information on the specifics of the study,
     there was sufficient informati
F007 Although a standard method was not followed, the testing procedures
     followed generally accepted fish bioconcentration guideline methods and
     although there was limited information on the specifics of the study,
     there was sufficient information on testing method and conditions in
     general to rate this study a 2.
F020 260545
EOR
F002 520
F010 3.7
F004 1
F005 RS
F006 At a concentration of 0.2 mg/L sec-butyl ether was shown to have
     bioconcentrate factor (BCF) range of 47 to 83, which is a log BCF range
     of 1.67 to 1.92.
* *
    At a concentration of 0.02 mg/L sec-butyl ether was shown to have a BCF
     range of 30
F007 At a concentration of 0.2 mg/L sec-butyl ether was shown to have
    bioconcentrate factor (BCF) range of 47 to 83, which is a log BCF range
    of 1.67 to 1.92.
* *
    At a concentration of 0.02 mg/L sec-butyl ether was shown to have a BCF
    range of 30 to 114, which is a log BCF range of 1.48 to 2.06.
F020 260543
EOR
F002 520
F010 3.7
F004 1
F005 TC
F006 Fish supplier was Sugishima fish farm, Kumamoto, Japan. Upon arrival,
     fish were placed in an acclimation tank with a flow through water system
     for 28 days at 25° +/-2° C. Test fish were then transferred to test tanks
     and reared again at the
F007 Fish supplier was Sugishima fish farm, Kumamoto, Japan. Upon arrival,
     fish were placed in an acclimation tank with a flow through water system
     for 28 days at 25^{\circ} +/-2° C. Test fish were then transferred to test tanks
     and reared again at the same temperature for approximately 1 month.
* *
* *
     At test initiation, average fish weight, length, and lipid content were,
     30 g, 10 cm, and 2-5%, respectively. Fish were fed pelleted feed for carp
     supplied by Haigo Shiryo K.K..
* *
* *
     Fish were fed approximately 2% of their total body weight daily. On days
     fish were sampled, they were not fed.
* *
     The test systems included 100 L volume glass tanks with a flow ate of 200
     to 800 ml/min at 25° +/-2° C. Dissolved oxygen was 6 to 8 mg/L. At test
     initiation there was a minimum of 15 fish per exposure level. Treatment
     levels were based on results of acute toxicity testing.
* *
```

Test water, test fish, and control fish were analyzed twice a week, every

two weeks, and prior to test initiation as well as at termination,

* *

```
respectively. Recovery efficiency was determined in water and fish
     homogenate. Analytical results from the definitive studies were corrected
     based on efficiency results.
* *
* *
     Bioconcentration factors were calculated based on:
* *
     test substance concentration in fish / test substance concentration in
     water.
* *
* *
     Two concentrations were evaluated: 0.2 mg/L and 0.02 mg/L.
* *
F020 260542
EOR
F002 520
F010 3.7
F004 1
F005 TS
F006 Analogue substance CAS No. 142-96-1; n-butyl ether
F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260541
EOR
F002 520
F010 3.7
F004 2
F005 ME
F006 Calculated values using BCFWIN version 2.15, a subroutine of the computer
    program EPIWIN version 3.12.
F007 Calculated values using BCFWIN version 2.15, a subroutine of the computer
     program EPIWIN version 3.12.
F020 260634
EOR
F002 520
F010 3.7
F004 2
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260637
EOR
F002 520
F010 3.7
F004 2
F005 RL
F006 The result is a calculated value based on the chemical structure and
     represents a potential bioaccumulation factor for the substance with the
     CAS number listed under test substance.
F007 The result is a calculated value based on the chemical structure and
     represents a potential bioaccumulation factor for the substance with the
     CAS number listed under test substance.
F020 260636
EOR
F002 520
F010 3.7
F004 2
F005 TC
```

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F006
* *
     BCFWIN estimates the bioconcentration factor (BCF) of an organic compound
     using the compound's log octanol-water partition coefficient (Kow).
* *
     The estimation methodology used by BCFWIN is described in "Improved
* *
     Method for Estimating Biocon
F007
     BCFWIN estimates the bioconcentration factor (BCF) of an organic compound
     using the compound's log octanol-water partition coefficient (Kow).
* *
     The estimation methodology used by BCFWIN is described in "Improved
     Method for Estimating Bioconcentration Factor (BCF) from Octanol-Water
     Partition Coefficient", SRC TR-97-006 (2nd Update), July 22, 1997.
* *
     Log Kow used = 2.87
* *
* *
* *
     Estimated BCF = 32.1
* *
     Estimated Log BCF = 1.507
* *
F020 260635
EOR
F002 520
F010 3.7
F004 2
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260633
EOR
F002 520
F010 4.1
F004 1
F005 CL
F006 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
     have an acute 96-hour LC50 of 14.7 mg/L and a Chronic Value of 2.2 mg/L
F007 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
     have an acute 96-hour LC50 of 14.7 mg/L and a Chronic Value of 2.2 mg/L
F020 260612
EOR
F002 520
F010 4.1
F004 1
F005 RE
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F020 260614
EOR
F002 520
F010 4.1
F004 1
F005 RL
F006 The value was calculated based on chemical structure as modeled by
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EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260613
EOR
F002 520
F010 4.1
F004 1
F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260611
EOR
F002 520
F010 4.1
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260554
EOR
F002 520
F010 4.1
F004 2
F005 CL
F006 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
     have an acute 96-hour LC50 of 10.8 mg/L and a Chronic Value of 1.6 mg/L.
F007 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
     have an acute 96-hour LC50 of 10.8 mg/L and a Chronic Value of 1.6 mg/L.
F020 260617
EOR
F002 520
F010 4.1
F004 2
F005 RE
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
```

```
EOR
F002 520
F010 4.1
F004 2
F005 RL
F006 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260618
EOR
F002 520
F010 4.1
F004 2
F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
    model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
* *
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260616
EOR
F002 520
F010 4.1
F004 2
F005 TS
F006 Analogue substance CAS No. 142-96-1; n-butyl ether
F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260615
EOR
F002 520
F010 4.1
F004 3
F006 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
     CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
    Ministry of Internatio
F007 Chemicals Inspection and Testing Institute (CITI) (1992). Biodegradation
     and bioaccumulation data of existing chemicals based on the CSCL Japan.
     CITI (ed.). Chemical Products Safety Division, Basic Industries Bureau,
    Ministry of International Trade and Industry, Japan. Japan Chemical
     Industry Ecology-Toxicology and Information Center.
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F020 260619

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F020 260622
EOR
F002 520
F010 4.1
F004 3
F005 RL
F006 Although a standard method was not followed, the testing procedures
     followed generally accepted fish acute toxicity guideline methods and
     although there was limited information on the specifics of the study,
     there was sufficient information
F007 Although a standard method was not followed, the testing procedures
     followed generally accepted fish acute toxicity guideline methods and
     although there was limited information on the specifics of the study,
     there was sufficient information on testing method and conditions in
     general to rate this study a 2.
F020 260623
EOR
F002 520
F010 4.1
F004 3
F005 TC
F006 Fish supplier was Nakashima fish farm, Kumamoto, Japan. Upon arrival,
     fish were placed in an acclimation tank with a flow-through water system
     for 3 to 5 weeks after external disinfection. The external disinfection
     was carried out under s
F007 Fish supplier was Nakashima fish farm, Kumamoto, Japan. Upon arrival,
     fish were placed in an acclimation tank with a flow-through water system
     for 3 to 5 weeks after external disinfection. The external disinfection
     was carried out under static conditions for about 24-hours in an acqueous
     solution containing 20 mg/l of Elbarju (Ueno Pharm. Co.) and 7 g/l sodium
     chloride.
* *
    Test fish were then placed in an acclimation tank with a flow-through
     system at 25°±2°C for about 28 days.
* *
    Dilution water for the test and culture was provided by a well of the
    Kurume Research Laboratories. Water temperature, pH, and dissolved
    oxygen were continuously monitored in the laboratory. Total hardness,
     evaporated residue, chemical oxygen demand, chloride ion, and other
    harmful substances were also monitored. The quality of the dilution
     water was confirmed to meet the standards of the Ministry of Health and
    Welfare in total hardness and evaporated residue. The other analyzed
     items met the water quality standards for fisheries.
* *
* *
    Test tanks were round glass vessels with 4 liters of test water per
     level. The test was conducted at 25^{\circ}\pm2^{\circ}C. Ten fish were tested at each
     level for 48 hours under semi-static conditions (renewal of test water
     every 8 to 16 hours).
* *
* *
    Test concentrations (levels) were not reported.
* *
* *
    The 48-hour LC50 value was estimated by either the Doudoroff Method or
     the Probit Method.
F020 260621
EOR
F002 520
F010 4.1
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```
F004 3
F005 TS
F006 Analogue substance CAS No. 142-96-1; n-butyl ether
F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260620
EOR
F002 520
F010 4.2
F004 1
F005 CL
F006 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
    have an acute 48-hour EC50 of 16.7 mg/L and a Chronic Value of 1.3 mg/L.
F007 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
    have an acute 48-hour EC50 of 16.7 mg/L and a Chronic Value of 1.3 mg/L.
F020 260625
EOR
F002 520
F010 4.2
F004 1
F005 RE
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F020 260627
EOR
F002 520
F010 4.2
F004 1
F005 RL
F006 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260626
EOR
F002 520
F010 4.2
F004 1
F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
* *
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
```

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* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260624
EOR
F002 520
F010 4.2
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260555
EOR
F002 520
F010 4.2
F004 2
F005 CL
F006 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
    have an acute 48-hour EC50 of 12.5 mg/L and a Chronic Value of 1.0 mg/L.
F007 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
    have an acute 48-hour EC50 of 12.5 mg/L and a Chronic Value of 1.0 mg/L.
F020 260630
EOR
F002 520
F010 4.2
F004 2
F005 RE
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F020 260632
EOR
F002 520
F010 4.2
F004 2
F005 RL
F006 The value was calculated based on chemical structure as modeled by
             This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260631
EOR
F002 520
F010 4.2
F004 2
F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
    model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
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model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260629
EOR
F002 520
F010 4.2
F004 2
F005 TS
F006 Analogue substance CAS No. 142-96-1; n-butyl ether
F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260628
EOR
F002 520
F010 4.3
F004 1
F005 CT
F006 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
    have an acute 96-hour EC50 of 11.0 mg/L and a Chronic Value of 1.8 mg/L.
F007 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
    have an acute 96-hour EC50 of 11.0 mg/L and a Chronic Value of 1.8 mg/L.
F020 260639
EOR
F002 520
F010 4.3
F004 1
F005 RE
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
    v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
F020 260641
EOR
F002 520
F010 4.3
F004 1
F005 RL
F006 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260640
EOR
F002 520
F010 4.3
F004 1
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F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
    model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260638
EOR
F002 520
F010 4.3
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260556
EOR
F002 520
F010 4.3
F004 2
F005 CL
F006 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
    have an acute 96-hour EC50 of 8.3 mg/L and a Chronic Value of 1.5 mg/L.
F007 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
    have an acute 96-hour EC50 of 8.3 mg/L and a Chronic Value of 1.5 mg/L.
F020 260644
EOR
F002 520
F010 4.3
F004 2
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F020 260646
EOR
F002 520
F010 4.3
F004 2
F005 RL
F006 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
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EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260645
EOR
F002 520
F010 4.3
F004 2
F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
    model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260643
EOR
F002 520
F010 4.3
F004 2
F005 TS
F006 Analogue substance CAS No. 142-96-1; n-butyl ether
F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260642
EOR
F002 520
F010 4.5.1
F004 1
F005 CL
F006 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
    have an acute 96-hour LC50 of 14.7 mg/L and a Chronic Value of 2.2 mg/L
F007 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
    have an acute 96-hour LC50 of 14.7 mg/L and a Chronic Value of 2.2 mg/L
F020 260649
EOR
F002 520
F010 4.5.1
F004 1
F005 RE
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F020 260651
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F002 520
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F010 4.5.1
F004 1
F005 RL
F006 The value was calculated based on chemical structure as modeled by
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     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260650
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F002 520
F010 4.5.1
F004 1
F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260648
EOR
F002 520
F010 4.5.1
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260647
EOR
F002 520
F010 4.5.1
F004 2
F005 CL
F006 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
     have an acute 96-hour LC50 of 10.8 mg/L and a Chronic Value of 1.6 mg/L.
F007 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
     have an acute 96-hour LC50 of 10.8 mg/L and a Chronic Value of 1.6 mg/L.
F020 260654
EOR
F002 520
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F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
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Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F020 260656
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F002 520
F010 4.5.1
F004 2
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F006 The value was calculated based on chemical structure as modeled by
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F007 The value was calculated based on chemical structure as modeled by
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     data are calculated and not measured.
F020 260655
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F002 520
F010 4.5.1
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F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
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     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
F020 260653
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F010 4.5.1
F004 2
F005 TS
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F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260652
EOR
F002 520
F010 4.5.2
F004 1
F005 CL
F006 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
     have an acute 48-hour EC50 of 16.7 mg/L and a Chronic Value of 1.3 mg/L.
F007 Based on the calculated Kow value of 2.87, sec-Butyl Ether is expected to
     have an acute 48-hour EC50 of 16.7 mg/L and a Chronic Value of 1.3 mg/L.
F020 260659
EOR
F002 520
F010 4.5.2
F004 1
F005 RE
F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
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Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
     Division. Washington, DC, USA.
F020 260661
EOR
F002 520
F010 4.5.2
F004 1
F005 RL
F006 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260660
EOR
F002 520
F010 4.5.2
F004 1
F005 TC
F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
    model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260658
EOR
F002 520
F010 4.5.2
F004 1
F005 TS
F006 CAS No. 6863-58-7; sec-butyl ether
F007 CAS No. 6863-58-7; sec-butyl ether
F020 260657
EOR
F002 520
F010 4.5.2
F004 2
F005 CL
F006 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
    have an acute 48-hour EC50 of 12.5 mg/L and a Chronic Value of 1.0 mg/L.
F007 Based on the calculated Kow value of 3.01, n-Butyl Ether is expected to
    have an acute 48-hour EC50 of 12.5 mg/L and a Chronic Value of 1.0 mg/L.
F020 260667
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EOR
F002 520
F010 4.5.2
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F006 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
F007 Cash G and Nabholz V (1999). ECOSAR Classes for Microsoft Windows, ECOWIN
     v0.99e. U.S. Environmental Protection Agency, OPPT - Risk Assessment
    Division. Washington, DC, USA.
F020 260669
EOR
F002 520
F010 4.5.2
F004 2
F005 RL
F006 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 260668
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F002 520
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F006 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
     model. The Kow calculation is performed by KOWWIN based on an
     atom/fragment contribution method of Meyl
F007 Log Kow (octanol/water partition coefficient) values and a chemical
     structure are needed to calculate aquatic toxicity using the ECOSAR
    model. The Kow calculation is performed by KOWWIN based on an
    atom/fragment contribution method of Meylan and Howard (1), which is a
     subroutine in the EPISuite computer model (2).
* *
* *
     1. Meylan, W. and P. Howard. 1995. Atom/fragment contribution method for
     estimating octanol-water partition coefficients. J. Pharm. Sci. 84:83-92.
* *
     2. EPI SuiteTM (2000). Estimation Program Interface Suite, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 260666
EOR
F002 520
F010 4.5.2
F004 2
F005 TS
F006 Analogue substance CAS No. 142-96-1; n-butyl ether
F007 Analogue substance CAS No. 142-96-1; n-butyl ether
F020 260665
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F005 1
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 2
F005 2
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 3
F005 3
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 4
F005 4
F006 27-10-2005
F007 10-02-2000
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EOR
F001 455
F002 1
F003 1.2
F004 5
F005 5
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 6
F005 6
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 8
F005 8
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 9
F005 9
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 10
F005 10
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 11
F005 11
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 1.2
F004 12
F005 12
F006 27-10-2005
F007 10-02-2000
EOR
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F001 455
F002 1
F003 1.2
F004 14
F005 14
F006 27-10-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 4.1
F004 1
F005 1
F006 01-11-2005
F007 10-02-2000
EOR
F001 455
F002 1
F003 4.2
F004 1
F005 1
F006 07-12-2005
F007 10-02-2000
EOB
C
B053 DS_REC_MARK_TAB
F001 455
F002 2.1
F003 2
F004 A37-009
EOR
F001 455
F002 2.2
F003 2
F004 A37-009
EOR
F001 455
F002 2.3
F003 2
F004 A37-009
EOR
F001 455
F002 2.4
F003 2
F004 A37-009
EOR
F001 455
F002 2.5
F003 2
F004 A37-009
EOR
F001 455
F002 2.6.1
F003 2
F004 A37-009
EOR
F001 455
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F002 3.1.1
F003 2
F004 A37-009
EOR
F001 455
F002 3.1.1
F003 3
F004 A37-009
EOR
F001 455
F002 3.1.2
F003 1
F004 A37-009
EOR
F001 455
F002 3.3.1
F003 2
F004 A37-009
EOR
F001 455
F002 3.3.1
F003 3
F004 A37-009
EOR
F001 455
F002 3.7
F003 1
F004 A37-009
EOR
F001 455
F002 4.1
F003 1
F004 A37-009
EOB
B051 DS_COMPONENT_TAB
F001 455
F002 0
F003 108-20-3
F012 N
F010 18-05-2005
F004 12032693
F005 18-05-2005
F006 12032693
F007 18-05-2005
F008 HPV
F009 A35-02
EOR
F001 455
F002 1
F003 108-20-3
F012 N
F010 10-02-2000
F011 10-02-2000
F004 101
F005 10-02-2000
F006 101
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F007 10-02-2000
F009 A35-02
EOB
B101 GI_GENERAL_INFORM_TAB
F001 455
F002 7
F003 27-10-2005
F004 CLGETTS
F010 A04-04
F011 A19-02
EOB
B102 GI_SYNONYM_TAB
F001 455
F002 1
F003 27-10-2005
F004 CLGETTS
F007 2,2'-oxybis-propane
EOR
F001 455
F002 2
F003 27-10-2005
F004 CLGETTS
F007 IPE
EOR
F001 455
F002 3
F003 27-10-2005
F004 CLGETTS
F007 2-isopropoxypropane
EOR
F001 455
F002 4
F003 27-10-2005
F004 CLGETTS
F007 IPE; Diisopropylether; DIPE; 2-Isopropoxy propane
EOR
F001 455
F002 5
F003 27-10-2005
F004 CLGETTS
F007 Diisopropyl Ether
EOR
F001 455
F002 6
F003 27-10-2005
F004 CLGETTS
F007 Isopropyl Ether
EOR
F001 455
F002 8
F003 27-10-2005
F004 CLGETTS
F007 2-Isopropoxy Propane
EOR
F001 455
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F002 9
F003 27-10-2005
F004 CLGETTS
F007 propane, 2,2'-oxybis-
EOR
F001 455
F002 10
F003 27-10-2005
F004 CLGETTS
F007 2,2'-oxybispropane
EOR
F001 455
F002 11
F003 27-10-2005
F004 CLGETTS
F007 Isopropylether
EOR
F001 455
F002 12
F003 27-10-2005
F004 CLGETTS
F007 Dipropyloxid
EOR
F001 455
F002 14
F003 27-10-2005
F004 CLGETTS
F007 2-Isopropoxypropan
EOB
B201 PC_MELTING_TAB
F001 455
F002 2
F003 07-12-2005
F004 CLGETTS
F015 A36-003
F016 1
F007 A02-03
F008 -86.8
F012 P01-03:
             not specified
F014 A03-02
F020 A01-03: Diisopropyl Ether (CAS # 108-20-3)
EOB
B202 PC_BOILING_TAB
F001 455
F002 2
F003 27-10-2005
F004 CLGETTS
F016 A36-003
F017 1
F007 A02-03
F008 68.5
F010 1013
F011 P02-01
F013 P03-03: not specified
F015 A03-02
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F018 A01-03: Diisopropylether
EOB
B203 PC_DENSITY_TAB
F001 455
F002 2
F003 07-12-2005
F004 CLGETTS
F016 A36-003
F017 1
F007 P05-02
F008 A02-03
F009 .7241
F011 P18-01
F012 20
F013 P04-03:
             not specified
F015 A03-02
F018 A01-03: Diisopropyl Ether (CAS # 108-20-3)
EOB
B204 PC_VAPOUR_TAB
F001 455
F002 2
F003 07-12-2005
F004 CLGETTS
F015 A36-003
F016 1
F007 A02-03
F008 198.65
F010 P02-01
F011 25
F014 A03-02
F018 A01-03: Diisopropyl Ether (CAS # 108-20-3)
B205 PC_PARTITION_TAB
F001 455
F002 2
F003 27-10-2005
F004 CLGETTS
F014 A36-003
F015 1
F007 A02-03
F008 1.52
F010 25
F011 P07-05
F013 A03-02
F016 A01-03: Diisopropylether
F020 C15-001
EOR
F001 455
F002 3
F003 12-12-2005
F004 CLGETTS
F014 A36-002
F015 2
F007 A02-03
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F008 2.4
F011 P07-04: Indirect method by reverse-phase HPLC
F013 A03-01
F016 A01-03: diisopropyl ether (CAS No. 108-20-3)
F020 C15-001
F019 6.7
EOB
B206 PC_WATER_SOL_TAB
F001 455
F002 2
F003 07-12-2005
F004 CLGETTS
F023 A36-003
F024 1
F007 A02-03
F008 P08-02
F009 8800
F011 20
F022 A03-02
             Diisopropyl Ether (CAS # 108-20-3)
F025 A01-03:
F030 C14-001
EOB
B301 EN_PHOTODEGRADATION_TAB
F001 455
F002 2
F003 07-12-2005
F004 CLGETTS
F045 A36-003
F046 1
F007 A01-03: Diisopropyl Ether (CAS # 108-20-3)
F008 F01-01
F009 F02-05: Calculated values using AOPWIN version 1.89, a subroutine of the
    computer program EPIWIN version 3.12
F023 25
F034 F06-03
F035 1500000
F036 F07-02
F044 A02-03
F037 .0000000002434
F038 A02-03
F040 50
F041 5.3
F042 F05-02
EOR
F001 455
F002 3
F003 07-12-2005
F004 CLGETTS
F045 A36-003
F046 2
F007 A01-03: Diisopropyl Ether (CAS # 108-20-3)
EOB
B302 EN_STABILITY_IN_WATER_TAB
F001 455
```

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F002 1
F003 07-12-2005
F004 CLGETTS
F040 A36-003
F041 1
F007 A01-03:
             Diisopropyl Ether (CAS # 108-20-3)
F008 F08-01
F009 F09-03: Technical discussion
F039 A03-02
EOB
B305 EN_TRANSPORT_TAB
F001 455
F002 2
F003 07-12-2005
F004 CLGETTS
F011 A36-003
F012 1
F008 F22-01: air - biota - sediment(s) - soil - water
F009 F21-01: Calculation according Mackay, Level I
EOR
F001 455
F002 3
F003 01-11-2005
F004 CLGETTS
F011 A36-003
F012 2
F007 F20-07
F008 F22-01
F009 F21-01: Level III simulation using the Mackay Multimedia Environmental
     Model (Mackay, 2001)
EOB
B308 EN_BIODEGRADATION_TAB
F001 455
F002 2
F003 07-12-2005
F004 CLGETTS
F047 A36-002
F048 1
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 F25-01
F009 F26-18
F010 1982
F011 F27-0139
F020 F30-02: not readily biodegradable
F046 A03-01
F052 28
F053 F05-01
EOR
F001 455
F002 3
F003 07-02-2006
F004 CLGETTS
F047 A36-003
F048 2
F007 A01-03: diisopropyl ether (CAS No. 108-20-3)
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F009 F26-25: American Public Health Association; No. 219 5-Day BOD; Standard
    Dilution Method
F010 1971
F011 F27-0166: sanitary waste treatment plant effluent
F046 A03-01
F052 5
F053 F05-01
EOR
F001 455
F002 4
F003 07-02-2006
F004 CLGETTS
F047 A36-003
F048 3
F007 A01-03:
              diisopropyl ether (CAS No. 108-20-3)
F008 F25-01
F009 F26-25: (comparison study of three aerobic biodegradation methods)
F010 1997
F011 F27-0137
F046 A03-01
EOR
F001 455
F002 5
F003 07-02-2006
F004 CLGETTS
F047 A36-003
F048 4
F007 A01-03: diisopropyl ether (CAS No. 108-20-3)
F008 F25-01
F009 F26-25: (continuous-flow bioreactors)
F010 2001
F011 F27-0166: Mixture (see remarks)
F046 A03-01
F052 600
F053 F05-01
EOR
F001 455
F002 6
F003 07-02-2006
F004 CLGETTS
F047 A36-003
F048 5
F007 A01-03: diisopropyl ether (CAS No. 108-20-3)
F009 F26-25: (soil/water microcosm)
F010 1999
F011 F27-0166: soil and groundwater from a site previously exposed to methyl
     tert-butyl ether
F046 A03-01
F052 1
F053 F05-05
EOR
F001 455
F002 7
F003 07-02-2006
F004 CLGETTS
F047 A36-003
F048 6
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F007 A01-03: diisopropyl ether (CAS No. 108-20-3)
F008 F25-02
F009 F26-25: (closed serum bottle test)
F010 1993
F011 F27-0166: sediment and groundwater from an anoxic aquifer polluted by
    municipal landfill leachate
F046 A03-01
F052 252
F053 F05-01
EOR
F001 455
F002 8
F003 24-02-2006
F004 CLGETTS
F047 A36-003
F048 7
F007 A01-03: diisopropyl ether (CAS No. 108-20-3)
F008 F25-02
F009 F26-25: unknown
F010 1994
F011 F27-0166: Sediment and surface or groundwater
F046 A03-01
EOR
F001 455
F002 9
F003 07-02-2006
F004 CLGETTS
F047 A36-003
F048 8
F007 A01-03: diisopropyl ether (CAS No. 108-20-3)
F008 F25-01
F009 F26-25: (sealed flasks, shaken)
F010 2000
F011 F27-0166: Gordonia terrae strain IFP 2001 (CNCM Registration No. CTP
     1-1889); isolated from activated sludge taken at an urban waste water
     treatment plant
F046 A03-01
F052 24
F053 F05-02
EOB
B310 EN_BIOACCUMULATION_TAB
F001 455
F002 1
F003 12-12-2005
F004 CLGETTS
F021 A36-003
F022 1
F007 A01-03: Diisopropyl Ether (CAS # 108-20-3)
F008 E02-0161: see remark
F009 F34-06: calculation
F015 25
F016 A02-03
F017 2.95
F020 A03-01
EOR
F001 455
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F002 2
F003 12-12-2005
F004 CLGETTS
F021 A36-003
F022 2
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 E02-0161: see remark
F009 F34-06: calculation
F015 25
F016 A02-03
F017 14.06
F020 A03-01
EOB
B311 EN_OTHER_TAB
F001 455
F002 1
F003 07-02-2006
F004 CLGETTS
F009 Biodegradation of diisopropyl ether
EOR
F001 455
F002 2
F003 27-02-2006
F004 CLGETTS
F010 2
F009 Biodegradation of diisopropyl ether under aerobic and anaerobic
     conditions - summary
EOB
B401 EC_FISHTOX_TAB
F001 455
F002 1
F003 01-11-2005
F004 CLGETTS
F033 A36-003
F034 1
F007 A01-03:
             Diisopropyl Ether (CAS No. 108-20-3)
F008 E01-02
F009 E02-0119
F010 E03-05: Flow-through Fish Acute Toxicity Test
F011 1983
F012 96
F013 E04-02
F014 E05-02
F021 A02-03
F022 91.7
F031 A03-03
F032 A03-02
EOR
F001 455
F002 2
F003 28-10-2005
F004 CLGETTS
F033 A36-003
F034 2
```

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F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F009 E02-0161: Fish
F010 E03-05: ECOSAR version 0.99h, US EPA
F012 96
F013 E04-02
F014 E05-02
F021 A02-03
F022 214.1
EOR
F001 455
F002 3
F003 12-12-2005
F004 CLGETTS
F033 A36-002
F034 3
F007 A01-03:
             Diisopropyl Ether (CAS No. 108-20-3)
F008 E01-02
F009 E02-0119
F010 E03-05: Flow-through Fish Acute Toxicity Test
F011 1983
F012 96
F013 E04-02
F014 E05-02
F021 A02-03
F022 786
F027 EC50
F028 A02-03
F029 476
F031 A03-03
F032 A03-02
EOR
F001 455
F002 4
F003 12-12-2005
F004 CLGETTS
F033 A36-002
F034 4
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 E01-02
F009 E02-0119
F010 E03-05: Flow-through Fish Acute Toxicity Test (ASTM, 1980)
F011 1985
F012 96
F013 E04-02
F014 E05-02
F021 A02-03
F022 900
F031 A03-03
F032 A03-02
EOR
F001 455
F002 5
F003 12-12-2005
F004 CLGETTS
F033 A36-004
F034 5
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
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F008 E01-05
F009 E02-0015
F010 E03-05: static acute fish toxicity test (APHA, 1971)
F012 24
F013 E04-02
F014 E05-02
F021 A02-03
F022 380
F031 A03-03
F032 A03-02
EOR
F001 455
F002 6
F003 12-12-2005
F004 CLGETTS
F033 A36-004
F034 6
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 E01-05
F009 E02-0070
F010 E03-05: static acute fish toxicity test
F012 96
F013 E04-02
F014 E05-02
F022 7000
F031 A03-01
F032 A03-01
EOR
F001 455
F002 7
F003 12-12-2005
F004 CLGETTS
F033 A36-004
F034 7
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 E01-05
F009 E02-0082
F010 E03-05: static acute fish toxicity test
F012 96
F013 E04-02
F014 E05-02
F022 6600
F031 A03-01
F032 A03-01
EOB
B402 EC_DAPHNIATOX_TAB
F001 455
F002 1
F003 07-12-2005
F004 CLGETTS
F032 A36-003
F033 1
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 E06-0010
F009 E07-04: U.S. Environmental Protection Agency, Methods for acute toxicity
     testing with fish, macro-invertebrates and amphibians (EPA-660/3-75-009)
```

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F010 1975
F011 48
F012 E04-02
F013 E05-02
F020 A02-03
F021 190
F030 A03-01
F031 A03-01
EOR
F001 455
F002 2
F003 28-10-2005
F004 CLGETTS
F032 A36-003
F033 2
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 E06-0034: Daphnia
F009 E07-04: ECOSAR version 0.99h, US EPA
F011 48
F012 E04-02
F013 E05-02
F020 A02-03
F021 221.9
EOB
B403 EC_ALGAETOX_TAB
F001 455
F002 2
F003 28-10-2005
F004 CLGETTS
F036 A36-003
F037 1
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 E08-0063: Green Alga
F009 E09-04: ECOSAR version 0.99h, US EPA
F012 96
F013 E04-02
F014 E05-02
F027 A02-03
F028 134.9
F030 ChV
F031 A02-03
F032 10.2
EOR
F001 455
F002 3
F003 12-12-2005
F004 CLGETTS
F036 A36-004
F037 2
F007 A01-03:
             Diisopropyl Ether (CAS No. 108-20-3)
F008 E08-0056
F009 E09-04: algae growth inhibition
F010 1983
F011 E10-01
F012 96
F013 E04-02
```

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F014 E05-02
F027 A02-05
F028 1000
F034 A03-01
F035 A03-02
EOB
B501 TO_ACUTE_ORAL_TAB
F001 455
F002 5
F003 01-11-2005
F004 CLGETTS
F017 A36-003
F018 1
F007 A01-03:
             Diisopropyl ether (CAS No. 108-20-3)
F008 T01-03
F009 T02-24
F010 T03-03: Similar to OECD 401
F016 A03-01
F019 T24-03
F021 T52-003: None; administered undiluted
F022 T23-42
EOR
F001 455
F002 6
F003 01-11-2005
F004 CLGETTS
F017 A36-003
F018 2
F007 A01-03: Diisopropyl ether (CAS No. 108-20-3)
F009 T02-23
F010 T03-03: Similar to OECD 401
F016 A03-01
F019 T24-04
F020 6
F021 T52-003: none reported
F022 T23-31
F023 8.2, 7.2, 6.0, 5.2, 3.3, 1.62 g/kg
EOB
B502 TO_ACUTE_INHAL_TAB
F001 455
F002 2
F003 01-11-2005
F004 CLGETTS
F019 A36-003
F020 1
F007 A01-03:
             Diisopropyl ether (CAS No. 108-20-3)
F009 T02-10
F010 T06-03: not specified
F018 A03-01
F021 T24-04
F023 T52-003: none
F024 T23-48: not specified
F025 0.3%; 1%; 3%; 6% in air
EOR
F001 455
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F002 3
F003 01-11-2005
F004 CLGETTS
F019 A36-003
F020 2
F007 A01-03: Diisopropyl ether (CAS No. 108-20-3)
F009 T02-23
F010 T06-03: not specified
F018 A03-01
F021 T24-04
F023 T52-003: none
F024 T23-31
F025 0.3%; 1%; 3%; 6% in air
EOR
F001 455
F002 4
F003 01-11-2005
F004 CLGETTS
F019 A36-003
F020 3
F007 A01-03:
             Diisopropyl ether (CAS No. 108-20-3)
F009 T02-17
F010 T06-03: not specified
F018 A03-01
F021 T24-01
F023 T52-003: none
F024 T23-48: Macacus rhesus
F025 0.3%; 1%; 3%; 6% in air
EOB
B503 TO_ACUTE_DERMAL_TAB
F001 455
F002 2
F003 01-11-2005
F004 CLGETTS
F017 A36-003
F018 1
F007 A01-03: Diisopropyl ether (CAS No. 108-20-3)
F008 T01-03
F009 T02-23
F010 T09-02: Similar to OECD 402
F016 A03-01
F019 T24-04
F021 T52-003: none
F022 T23-31
F023 variable
EOB
B507 TO_SENSITIZATION_TAB
F001 455
F002 2
F003 01-11-2005
F004 CLGETTS
F015 A36-003
F016 1
F007 A01-03: Diisopropyl ether (CAS No.108-20-3)
F008 T18-14: In vitro chemical reactivity assay, surrogate for respiratory
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* sensitization
F009 T02-19: No animals; in vitro chemical assay
F010 T20-03: No guideline available
F011 1990
F012 T47-01
F013 T21-02
F014 A03-01
F017 0
F030 T52-003: None
EOB
B508 TO_REPEATED_DOSE_TAB
F001 455
F002 5
F003 01-11-2005
F004 CLGETTS
F030 A36-003
F031 1
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 T02-24
F009 T23-42
F010 T24-03
F011 T25-08
F012 T26-23
F013 1996
F014 6 hours/day
F015 5 days/week for ~13 weeks
F017 0, 480, 3300, or 7100 ppm
F018 T27-03: yes (untreated & sham-exposed)
F019 A02-03
F020 480
F022 T28-05
F029 A03-02
F032 C07-002
EOR
F001 455
F002 6
F003 01-11-2005
F004 CLGETTS
F030 A36-003
F031 2
F007 A01-03:
             Diisopropyl Ether (CAS No. 108-20-3)
F008 T02-24
F009 T23-42
F010 T24-03
F011 T25-08
F012 T26-16: USEPA 1984, 40CFR Part 798:6050, 6400, and 6200
F013 1997
F014 6 hours/day
F015 5 days/week for ~13 weeks
F017 0, 450, 3250, or 7060 ppm
F018 T27-03: yes (sham-exposed)
F029 A03-02
F032 C07-002
EOB
B509 TO_GENETIC_IN_VITRO_TAB
```

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F001 455
F002 4
F003 10-11-2005
F004 CLGETTS
F016 A36-003
F017 1
F007 A01-03:
             Diisopropyl ether (CAS No. 108-20-3)
F008 T30-05
F009 T31-18: Similar to OECD Guideline 471
F010 1988
F011 Salmonella typhimurium
F012 T32-03
F013 T33-02
F014 A03-02
F015 Up to 8000 ug/ml in the pre-incubation mix
EOR
F001 455
F002 5
F003 10-11-2005
F004 CLGETTS
F016 A36-003
F017 2
F007 A01-03: Di-isopropyl ether (CAS No. 108-20-3)
F008 T30-15
F009 T31-18: Similar to OECD Guideline 473
F010 1984
F011 Chinese hamster ovary cells
F012 T32-04
F013 T33-02
F014 A03-02
F015 Up to 1200 ug/ml
EOR
F001 455
F002 6
F003 10-11-2005
F004 CLGETTS
F016 A36-003
F017 3
F007 A01-03: Di-isopropyl ether (CAS No. 108-20-3)
F008 T30-07
F009 T31-18: Similar to OECD Guideline 476
F010 1984
F011 Rat liver cells
F012 T32-04
F013 T33-02
F014 A03-02
F015 Up to 1200 ug/ml
EOR
F001 455
F002 7
F003 10-11-2005
F004 CLGETTS
F016 A36-003
F017 4
F007 A01-03:
              Di-isopropyl ether (CAS No. 108-20-3)
F008 T30-09
F009 T31-18: Similar to OECD Guideline 481
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```
F010 1984
F011 Saccharomyces cerevisiae
F012 T32-03
F013 T33-02
F014 A03-02
F015 Up to 8000 ug/ml in the pre-incubation mix
B513 TO_DEVELOPMENTAL_TAB
F001 455
F002 2
F003 01-11-2005
F004 CLGETTS
F030 A36-003
F007 A01-03: Diisopropyl Ether (CAS No. 108-20-3)
F008 T02-24
F009 T23-42
F010 T24-01
F011 T25-08
F012 T44-06
F013 1996
F014 20 days
F015 6 hr/day
F016 Gestation Days 6-15
F017 0, 430, 3095, or 6745 ppm
F018 T27-03: yes (untreated & sham-exposed)
F029 A03-02
F032 T58-007: NOEL Maternal
F033 A02-03
F034 430
F036 T43-04
F037 T58-007: NOEL Pup
F038 A02-03
F039 430
F041 T43-04
F047 Maternal NOEL: 430 ppm; Pup NOEL: 430 ppm
EOB
B514 TO_OTHER_TAB
F001 455
F002 2
F003 01-11-2005
F004 CLGETTS
F008 A36-003
F009 1
F007 T45-12: Sensory Irritation in Humans
EOR
F001 455
F002 3
F003 01-11-2005
F004 CLGETTS
F008 A36-003
F009 2
F007 T45-12: Sensory irritation in humans
EOB
B601 TEXT_TAB
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F002 455
F010 2.1
F004 2
F005 RE
F006 Lide D, et al. (eds.) (1997-1998). CRC Handbook of Chemistry and Physics.
     78th Edition. CRC Press, New York, NY, USA.
F007 Lide D, et al. (eds.) (1997-1998). CRC Handbook of Chemistry and Physics.
     78th Edition. CRC Press, New York, NY, USA.
F020 241178
EOR
F002 455
F010 2.1
F004 2
F005 RL
F006 The CRC Handbook of Chemistry and Physics is a peer reviewed publication.
     This robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F007 The CRC Handbook of Chemistry and Physics is a peer reviewed publication.
     This robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F020 241177
EOR
F002 455
F010 2.1
F004 2
F005 TS
F006 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F007 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F020 241176
EOR
F002 455
F010 2.2
F004 2
F005 RE
F006 Lide D, et al. (eds.) (1997-1998). CRC Handbook of Chemistry and Physics.
     78th Edition. CRC Press, New York, NY, USA.
F007 Lide D, et al. (eds.) (1997-1998). CRC Handbook of Chemistry and Physics.
     78th Edition. CRC Press, New York, NY, USA.
F020 241181
EOR
F002 455
F010 2.2
F004 2
F005 RL
F006 The CRC Handbook of Chemistry and Physics is a peer reviewed publication.
     This robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F007 The CRC Handbook of Chemistry and Physics is a peer reviewed publication.
     This robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F020 241180
EOR
F002 455
F010 2.2
F004 2
F005 TS
F006 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
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F007 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F020 241179
EOR
F002 455
F010 2.3
F004 2
F005 RE
F006 Lide D, et al. (eds.) (1997-1998). CRC Handbook of Chemistry and Physics.
     78th Edition. CRC Press, New York, NY, USA.
F007 Lide D, et al. (eds.) (1997-1998). CRC Handbook of Chemistry and Physics.
     78th Edition. CRC Press, New York, NY, USA.
F020 241184
EOR
F002 455
F010 2.3
F004 2
F005 RL
F006 The CRC Handbook of Chemistry and Physics is a peer reviewed publication.
     This robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F007 The CRC Handbook of Chemistry and Physics is a peer reviewed publication.
     This robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F020 241183
EOR
F002 455
F010 2.3
F004 2
F005 TS
F006 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F007 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F020 241182
EOR
F002 455
F010 2.4
F004 2
F005 ME
F006 Method not specified.
F007 Method not specified.
F020 241243
EOR
F002 455
F010 2.4
F004 2
F005 RE
F006 Daubert T and Danner R (1989). Physical and thermodynamic properties of
    pure chemicals: Data compilation. Design Institute for Physical Property
    Data, American Institute of Chemical Engineers. Hemisphere Publishing
     Corp., New York, NY, USA.
F007 Daubert T and Danner R (1989). Physical and thermodynamic properties of
     pure chemicals: Data compilation. Design Institute for Physical Property
     Data, American Institute of Chemical Engineers. Hemisphere Publishing
     Corp., New York, NY, USA.
F020 241187
EOR
F002 455
F010 2.4
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F004 2
F005 RL
F006 This robust summary has a reliability rating of 2 because the data were
     not reviewed for quality, however, the reference is from a peer-reviewed
     handbook.
F007 This robust summary has a reliability rating of 2 because the data were
     not reviewed for quality, however, the reference is from a peer-reviewed
     handbook.
F020 241186
EOR
F002 455
F010 2.4
F004 2
F005 TS
F006 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F007 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F020 241185
EOR
F002 455
F010 2.5
F004 2
F005 ME
F006 Method not specified.
F007 Method not specified.
F020 241191
EOR
F002 455
F010 2.5
F004 2
F005 RE
F006 Hansch C, Leo A and Hoekman D (1995). Exploring QSAR - Hydrophobic,
     Electronic and Steric Constants. p. 6. ACS Professional Reference Book,
     American Chemical Society, Washington, DC, USA.
F007 Hansch C, Leo A and Hoekman D (1995). Exploring QSAR - Hydrophobic,
     Electronic and Steric Constants. p. 6. ACS Professional Reference Book,
     American Chemical Society, Washington, DC, USA.
F020 241194
EOR
F002 455
F010 2.5
F004 2
F005 RL
F006 The value cited by the authors is a measured and preferred value. This
     robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F007 The value cited by the authors is a measured and preferred value. This
     robust summary has a reliability rating of 2 because there is
     insufficient information available on the method and analytical procedure.
F020 241193
EOR
F002 455
F010 2.5
F004 2
F005 TS
F006 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F007 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F020 241192
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EOR
F002 455
F010 2.5
F004 3
F005 RE
F006 Eadsforth C (1983). Isopropyl ether: determination of the n-octanol/water
     partition coefficient using a reverse-phase HPLC method. Report #
     SBGR.83.131. Shell Research Limited, Sittingbourne Research Centre,
     Sittingbourne, Kent, England.
F007 Eadsforth C (1983). Isopropyl ether: determination of the n-octanol/water
     partition coefficient using a reverse-phase HPLC method. Report #
     SBGR.83.131. Shell Research Limited, Sittingbourne Research Centre,
     Sittingbourne, Kent, England.
F020 243466
EOR
F002 455
F010 2.5
F004 3
F005 RS
F006 Log Pow = 2.4 (Pow = 250) at pH 6.7
F007 \text{ Log Pow} = 2.4 \text{ (Pow} = 250) at pH 6.7
F020 243464
EOR
F002 455
F010 2.5
F004 3
F005 TC
F006 The HPLC system was a reverse-phase C18-coated silica gel column, 250 mm
     {\tt x} 5 mm id, with a mobile phase of 3 volumes methanol and 1 volume water
     (final pH 6.7) at a flow rate of 1 mL/min. 100 mL samples of an
     approximate 1 mg/mL solution i
F007 The HPLC system was a reverse-phase C18-coated silica gel column, 250 mm
     x 5 mm id, with a mobile phase of 3 volumes methanol and 1 volume water
     (final pH 6.7) at a flow rate of 1 mL/min. 100 mL samples of an
     approximate 1 mg/mL solution in the mobile phase were injected, and the
     emergence of the material was observed using refraction index detection.
     Thirty-one reference compounds were used to generate the linear
     relationship between log k (k = capacity factor) and log Pow. Using the
     \operatorname{HPLC} retention time for the peak of the test substance, the \log k was
     determined, and the log Pow value was calculated using the linear
     equation developed from the reference compounds.
* *
     Log Pow was determined according to the following calculations:
* *
     Retention time (RT), min = 5.7
* *
     Capacity factor, k = 0.87, k = (RTcmpd - RTunretained std)/RTunretained
     std
* *
     log k = -0.06
* *
     linear equation: log k = -0.930 + 0.357 log Pow
F020 243465
EOR
F002 455
F010 2.6.1
F004 2
F005 RE
F006 Gerhartz W, Yamamoto Y, Kaudy L, Rounsaville J and Schulz G (eds.)
     (1987). Ullmann's Encyclopedia of Industrial Chemistry. Vol. A 10. 5th
     Edition. VCH Publishers, New York, NY, USA.
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F007 Gerhartz W, Yamamoto Y, Kaudy L, Rounsaville J and Schulz G (eds.)
     (1987). Ullmann's Encyclopedia of Industrial Chemistry. Vol. A 10. 5th
     Edition. VCH Publishers, New York, NY, USA.
F020 241190
EOR
F002 455
F010 2.6.1
F004 2
F005 RL
F006 The Ullmann's Encyclopedia of Industrial Chemistry is a peer reviewed
     publication. This robust summary has a reliability rating of 2 because
     there is insufficient information available on the method and analytical
     procedure.
F007 The Ullmann's Encyclopedia of Industrial Chemistry is a peer reviewed
     publication. This robust summary has a reliability rating of 2 because
     there is insufficient information available on the method and analytical
     procedure.
F020 241189
EOR
F002 455
F010 2.6.1
F004 2
F005 TS
F006 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F007 CAS No. 108-20-3; diisopropyl ether; purity is unknown.
F020 241188
EOR
F002 455
F010 3.1.1
F004 2
F005 ME
F006 Calculated values using AOPWIN version 1.89, a subroutine of the computer
    program EPIWIN version 3.12
* *
* *
     Indirect photodegradation, or atmospheric oxidation potential, is based
     on the structure-activity relationship methods developed by R. At
F007 Calculated values using AOPWIN version 1.89, a subroutine of the computer
    program EPIWIN version 3.12
* *
     Indirect photodegradation, or atmospheric oxidation potential, is based
     on the structure-activity relationship methods developed by R. Atkinson
    under the following conditions:
* *
       Temperature: 25°C
* *
       Sensitizer: OH- radical
* *
       Concentration of Sensitizer: 1.5E6 OH- radicals/cm3
F020 241161
EOR
F002 455
F010 3.1.1
F004 2
F005 RE
F006 EPIWIN (2000). Estimation Program Interface for Windows, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPIWIN (2000). Estimation Program Interface for Windows, version 3.12.
     Syracuse Research Corporation, Syracuse, NY, USA.
F020 241164
EOR
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```
F002 455
F010 3.1.1
F004 2
F005 RL
F006 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F007 The value was calculated based on chemical structure as modeled by
     EPIWIN. This robust summary has a reliability rating of 2 because the
     data are calculated and not measured.
F020 241163
EOR
F002 455
F010 3.1.1
F004 2
F005 RM
F006 DIPE has the potential to volatilize to air, based on a vapor pressure of
     19,865 Pa at 25°C (Daubert and Danner, 1989), where it is subject to
     atmospheric oxidation. In air, DIPE can react with photosensitized oxygen
     in the form of hydroxyl
F007 DIPE has the potential to volatilize to air, based on a vapor pressure of
     19,865 Pa at 25°C (Daubert and Danner, 1989), where it is subject to
     atmospheric oxidation. In air, DIPE can react with photosensitized oxygen
     in the form of hydroxyl radicals (OH-). The computer program AOPWIN
     (atmospheric oxidation program for Microsoft Windows) (EPIWIN, 2000)
     calculates a chemical half-life for a 12-hour day (the 12-hour day
    half-life value normalizes degradation to a standard day light period
    during which hydroxyl radicals needed for degradation are generated),
    based on an OH- reaction rate constant and a defined OH- concentration.
    DIPE has a calculated half-life in air of 5.3 hours or 0.4 days (12-hour
     day), based on a rate constant of 24.34 E-12 cm3/molecule*sec and an OH-
     concentration of 1.5 E5 OH-/cm3.
F020 241162
EOR
F002 455
F010 3.1.1
F004 3
F005 ME
F006 Technical discussion
F007 Technical discussion
F020 241165
EOR
F002 455
F010 3.1.1
F004 3
F005 RE
F006 Zepp R and Cline D (1977). Rates of direct photolysis in the aqueous
     environment. Environ. Sci. Technol. 11:359-366.
F007 Zepp R and Cline D (1977). Rates of direct photolysis in the aqueous
     environment. Environ. Sci. Technol. 11:359-366.
F020 241167
EOR
F002 455
F010 3.1.1
F004 3
F005 RL
F006 This robust summary has a reliability of 2 because it is a technical
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discussion and not a study.
F007 This robust summary has a reliability of 2 because it is a technical
     discussion and not a study.
F020 243461
EOR
F002 455
F010 3.1.1
F004 3
F005 RM
F006 Direct photochemical degradation occurs through the absorbance of solar
     radiation by a chemical substance in aqueous solution. If the absorbed
     energy is high enough, then the resultant excited state of the chemical
     may undergo a transformat
F007 Direct photochemical degradation occurs through the absorbance of solar
     radiation by a chemical substance in aqueous solution. If the absorbed
     energy is high enough, then the resultant excited state of the chemical
     may undergo a transformation. A prerequisite for direct photodegradation
     is the ability of one or more bonds within a chemical to absorb
     ultraviolet (UV)/visible light in the 290 to 750 nm range. Light
     wavelengths longer than 750 nm do not contain sufficient energy to break
     chemical bonds, and wavelengths below 290 nm are shielded from the earth
    by the stratospheric ozone layer (Harris, 1982a).
    An approach to assessing the potential for a substance to undergo
    photochemical degradation is to assume that degradation will occur in
    proportion to the amount of light wavelengths >290 nm absorbed by
     constituent molecules (Zepp and Cline, 1977). The oxygen non-bonding
     electrons in ethers do not give rise to absorption above 160 nm, which is
     why pure ether solvents can be used in spectroscopic studies.
     Consequently, DIPE is not subject to photolytic processes in the aqueous
     environment.
F020 241166
EOR
F002 455
F010 3.1.2
F004 1
F005 RE
F006 Gould E (1959). Mechanism and Structure in Organic Chemistry. Holt,
    Reinhart and Winston, New York, NY, USA.
F007 Gould E (1959). Mechanism and Structure in Organic Chemistry. Holt,
    Reinhart and Winston, New York, NY, USA.
F020 241169
EOR
F002 455
F010 3.1.2
F004 1
F005 RE
F006 Harris J (1982). Rate of Hydrolysis. In: Handbook of Chemical Property
     Estimation Methods. Chapter 7. Edited by WJ Lyman, WF Reehl and DH
     Rosenblatt. McGraw-Hill Book Company, New York, NY, USA.
F007 Harris J (1982). Rate of Hydrolysis. In: Handbook of Chemical Property
     Estimation Methods. Chapter 7. Edited by WJ Lyman, WF Reehl and DH
     Rosenblatt. McGraw-Hill Book Company, New York, NY, USA.
F020 241170
EOR
F002 455
F010 3.1.2
F004 1
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F005 RL
F006 This robust summary has a reliability of 2 because it is a technical
     discussion and not a study.
F007 This robust summary has a reliability of 2 because it is a technical
     discussion and not a study.
F020 243463
EOR
F002 455
F010 3.1.2
F004 1
F005 RS
F006 Hydrolysis of an organic chemical is the transformation process in which
     a water molecule or hydroxide ion reacts to form a new carbon-oxygen
    bond. Chemicals with leaving groups that have a potential to hydrolyze
     include alkyl halides, amid
F007 Hydrolysis of an organic chemical is the transformation process in which
     a water molecule or hydroxide ion reacts to form a new carbon-oxygen
    bond. Chemicals with leaving groups that have a potential to hydrolyze
     include alkyl halides, amides, carbamates, carboxylic acid esters and
     lactones, epoxides, phosphate esters, and sulfonic acid esters (Gould,
     1959). The lack of a suitable leaving group renders a compound resistant
     to hydrolysis. DIPE is resistant to hydrolysis because it lacks a
     functional group that is hydrolytically reactive and Harris (1982b)
     identifies ether groups as generally resistant to hydrolysis. Therefore,
    hydrolysis will not contribute to the removal of diisopropyl ether from
     the environment.
F020 241168
EOR
F002 455
F010 3.3.1
F004 2
F005 RE
F006 Mackay D (1998). Level I Fugacity-Based Environmental Equilibrium
     Partitioning Model, Version 2.1 (16-bit). Environmental Modelling
     Centre, Trent University, Ontario, Canada.
F007 Mackay D (1998). Level I Fugacity-Based Environmental Equilibrium
     Partitioning Model, Version 2.1 (16-bit). Environmental Modelling
     Centre, Trent University, Ontario, Canada.
F020 241175
EOR
F002 455
F010 3.3.1
F004 2
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated.
F020 241174
EOR
F002 455
F010 3.3.1
F004 2
F005 RM
F006 Physicochemical data used in the calculation:
* *
* *
    Parameter
                       Value w/ Units
```

```
* *
     Molecular Weight = 102.18
* *
     Temperature = 25° C
* *
     Log Kow = 1.52
* *
     Water Solubility = 8,800 g/m3
* *
     Vapor Pressure = 19,865 Pa
* *
     Melting Point = -86.8^{\circ} C
F007 Physicochemical data used in the calculation:
* *
* *
     Parameter
                        Value w/ Units
* *
* *
     Molecular Weight = 102.18
* *
     Temperature = 25° C
     Log Kow = 1.52
* *
* *
     Water Solubility = 8,800 g/m3
* *
     Vapor Pressure = 19,865 Pa
* *
     Melting Point = -86.8^{\circ} C
F020 241171
EOR
F002 455
F010 3.3.1
F004 2
F005 RS
F006 Using the Mackay Level I calculation, the following
     distribution is predicted for diisopropyl ether:
* *
* *
     %Distribution
                       Compartment
* *
        97.83
                        Air
* *
         2.10
                        Water
* *
         0.06
                        Soil
* *
         <0.01
                        Sediment
* *
         <0.01
F007 Using the Mackay Level I calculation, the following
     distribution is predicted for diisopropyl ether:
* *
* *
     %Distribution
                       Compartment
* *
       97.83
                        Air
* *
         2.10
                        Water
         0.06
                        Soil
* *
                        Sediment
         <0.01
* *
         <0.01
                        Suspended Sediment
* *
         <0.01
                        Biota
F020 241172
EOR
F002 455
F010 3.3.1
F004 2
F005 TS
F006 Diisopropyl Ether (CAS # 108-20-3)
F007 Diisopropyl Ether (CAS # 108-20-3)
F020 241173
EOR
F002 455
F010 3.3.1
F004 3
F005 CL
F006 The majority of DIPE is calculated to partition into the water phase,
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with smaller but significant amounts into air and soil, based on the
     modeling parameters used in this calculation. DIPE is considered to be a
     Type 1 chemical with potenti
F007 The majority of DIPE is calculated to partition into the water phase,
     with smaller but significant amounts into air and soil, based on the
     modeling parameters used in this calculation. DIPE is considered to be a
     Type 1 chemical with potential to partition into all environmental
     compartments.
F020 241237
EOR
F002 455
F010 3.3.1
F004 3
F005 ME
F006 Level III simulation using the Mackay Multimedia Environmental Model
     (Mackay, 2001). Mass balances are calculated for the four bulk media of
     air (gas + aerosol), water (solution + suspended sediment + biota), soil,
     (solids + air + water), a
F007 Level III simulation using the Mackay Multimedia Environmental Model
     (Mackay, 2001). Mass balances are calculated for the four bulk media of
     air (gas + aerosol), water (solution + suspended sediment + biota), soil,
     (solids + air + water), and sediment (solids + pore water). Equilibrium
     exists within, but not between media. Physical-chemical properties are
    used to quantify a chemical's behavior in an evaluative environment.
    Three types of chemicals are treated in this model: chemicals that
    partition into all media (Type 1), non volatile chemicals (Type 2), and
     chemicals with zero, or near-zero, solubility (Type 3). The model can not
     treat ionizing or speciating substances. The Level III model assumes a
     simple, evaluative environment with user-defined volumes and densities
     for the following homogeneous environmental media (or compartments): air,
     water, soil, sediment, suspended sediment, fish and aerosols.
* *
    This model provides a description of a chemical's fate including the
     important degradation and advection losses and the intermedia transport
    processes. The distribution of the chemical between media depends on how
     the chemical enters the system, e.g. to air, to water, or to both. This
    mode of entry also affects persistence or residence time.
    The rates of intermedia transport are controlled by a series of 12
     transport velocities. Reaction half-lives are requested for all 7 media.
     The advective residence time selected for air also applies to aerosols
     and the residence time for water applies to suspended sediment and fish.
     The advective residence time of aerosols, suspended sediment and fish
     cannot be specified independently of the air and water residence times.
F020 241234
EOR
F002 455
F010 3.3.1
F004 3
F005 RE
F006 Mackay D (1991). Multimedia Environmental Models; The Fugacity Approach.
     Lewis Publishers, CRC Press, pp 67-183.
F007 Mackay D (1991). Multimedia Environmental Models; The Fugacity Approach.
     Lewis Publishers, CRC Press, pp 67-183.
F020 241239
EOR
F002 455
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F010 3.3.1
F004 3
F005 RE
F006 Mackay D (2001). Multimedia Environmental Models: The Fugacity Approach -
     Second Edition. Lewis Publishers, Boca Raton, pp.1-261.
F007 Mackay D (2001). Multimedia Environmental Models: The Fugacity Approach -
     Second Edition. Lewis Publishers, Boca Raton, pp.1-261.
F020 241242
EOR
F002 455
F010 3.3.1
F004 3
F005 RE
F006 Mackay D, et al (1996a). Assessing the fate of new and existing
     chemicals: a five-stage process. Environ. Toxicol. Chem. 15(9):1618-1626.
F007 Mackay D, et al (1996a). Assessing the fate of new and existing
     chemicals: a five-stage process. Environ. Toxicol. Chem. 15(9):1618-1626.
F020 241240
EOR
F002 455
F010 3.3.1
F004 3
F005 RE
F006 Mackay D, et al (1996b). Evaluating the environmental fate of a variety
     of types of chemicals using the EQC model. Environ. Toxicol. Chem.
     15(9):1627-1637.
F007 Mackay D, et al (1996b). Evaluating the environmental fate of a variety
     of types of chemicals using the EQC model. Environ. Toxicol. Chem.
     15(9):1627-1637.
F020 241241
EOR
F002 455
F010 3.3.1
F004 3
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated.
F020 241238
EOR
F002 455
F010 3.3.1
F004 3
F005 RS
F006 Output
* *
            Mass%
                         Half life(hr)
                                           Emissions(kg/hr)
* *
                  19.4
     Air
                               25.2
                                           1000
* *
                  61.0
                               360
                                           1000
     Water
* *
     Soil
                  19.5
                               720
                                           1000
* *
     Sediment
                   0.1
                               3240
F007 Output
* *
                         Half life(hr)
                                           Emissions(kg/hr)
            Mass%
* *
                  19.4
                               25.2
                                           1000
     Air
* *
     Water
                  61.0
                               360
                                           1000
* *
                  19.5
                               720
                                           1000
     Soil
* *
     Sediment
                   0.1
                               3240
                                               0
```

```
F020 241236
EOR
F002 455
F010 3.3.1
F004 3
F005 TC
F006 Physchem Inputs
     Molar Mass = 102.18
* *
     Data Temperature = 25 °C
* *
     Water Solubility = 8800 mg/l exp.
* *
     Vapour Pressure = 19865 Pa exp.
* *
     Log Kow = 1.52 \exp.
* *
     Melting Point = -86.8 °C exp.
* *
* *
     Reaction Half Lives in hours (if not available they can be pre
F007 Physchem Inputs
* *
     Molar Mass = 102.18
* *
     Data Temperature = 25 °C
* *
     Water Solubility = 8800 mg/l exp.
* *
     Vapour Pressure = 19865 Pa exp.
* *
     Log Kow = 1.52 \exp.
* *
     Melting Point = -86.8 °C exp.
* *
* *
     Reaction Half Lives in hours (if not available they can be predicted
     using EPIWIN)
* *
                                25.2
     Air (gaseous)
* *
     Water (no susp. part.)
                                360
* *
     Bulk Soil
     Bulk Sediment
                                3240
* *
     Suspended Particles
                                360
* *
     Fish
                         360
* *
                                      25.2
     Aerosol
* *
* *
     Environmental Properties (EQC standard environment)
* *
     Dimensions (all defaults)
* *
     Densities (all defaults)
* *
     Organic carbon & Advection (all defaults)
* *
     Transport Velocities (all defaults)
* *
     Emission and Inflows (defaults used)
* *
     Air 1000 kg/hr
* *
     Water 1000 kg/hr
* *
     Soil 1000 kg/hr
* *
     Sediment 0 kg/hr
F020 241235
EOR
F002 455
F010 3.3.1
F004 3
F005 TS
F006 Diisopropyl Ether, CAS No. 108-20-3
F007 Diisopropyl Ether, CAS No. 108-20-3
F020 241233
EOR
F002 455
F010 3.5
F004 2
```

```
F005 CL
F006 Diisopropyl ether is not readily biodegradable and it did not
     significantly inhibit the biodegradability of the test substance in an
     inhibition test.
F007 Diisopropyl ether is not readily biodegradable and it did not
     significantly inhibit the biodegradability of the test substance in an
     inhibition test.
F020 241218
EOR
F002 455
F010 3.5
F004 2
F005 RE
F006 Stone C and Watkinson R (1983). Isopropyl ether: An assessment of ready
     biodegradability. Report # SBGR.83.428. Shell Biosciences Laboratory,
     Sittingbourne Research Centre, Sittingbourne, Kent, England.
F007 Stone C and Watkinson R (1983). Isopropyl ether: An assessment of ready
     biodegradability. Report # SBGR.83.428. Shell Biosciences Laboratory,
     Sittingbourne Research Centre, Sittingbourne, Kent, England.
F020 241219
EOR
F002 455
F010 3.5
F004 2
F005 RS
F006 Test substance was not readily biodegradable. After 28 days, the test
     substance exhibited no measurable biodegradation. By day 5, >60%
     biodegradation of positive control was observed, which meets the
     guideline requirement. No excursions fro
F007 Test substance was not readily biodegradable. After 28 days, the test
     substance exhibited no measurable biodegradation. By day 5, >60%
     biodegradation of positive control was observed, which meets the
     guideline requirement. No excursions from the testing guideline were
     noted. The inhibition study showed that the test substance did not
     inhibit the biodegradability of the positive control substance, sodium
     benzoate.
* *
                                % Degradation* Mean
                                                             % Degradation
                        (day 28)
     Sample
                                                                 (day 28)
* *
* *
     Test Substance
                                0.0, 0.0
                                                                          0.0
                               65.0, 73.0
                                                                       69.0
* *
     Na Benzoate
* *
     * duplicate data
* *
* *
     Mean oxygen concentrations (mg/L) of duplicate test systems:
* *
     Day 0
* *
     Mineral Salts Control = 8.85
* *
     Blank = 8.8
* *
     Na Benzoate = 8.95
* *
     Test Substance = 8.9 (single test system)
* *
     Test Substance + Na Benzoate = 8.9* (single test system)
* *
     Day 5
* *
    Mineral Salts Control = 9.0
     Blank = 8.8
* *
* *
     Na Benzoate = 5.7
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```
* *
     Test Substance + Na Benzoate = 5.8
* *
* *
     Day 15
* *
     Mineral Salts Control = 8.75
* *
     Blank = 8.65
* *
     Na Benzoate = 4.9
* *
     Test Substance = 8.55
* *
     Test Substance + Na Benzoate = 4.9
* *
* *
     Day 28
* *
     Mineral Salts Control = 8.65
* *
     Blank = 7.05
* *
     Na Benzoate = 3.6
* *
     Test Substance = 8.3
* *
     Test Substance + Na Benzoate = 4.15
F020 241217
EOR
F002 455
F010 3.5
F004 2
F005 TC
F006 The inoculum source was the Sittingbourne Sewage works in Kent, England,
     and was prepared according to methods described in the OECD 301D
     guideline. The test substance was added to the test medium by direct
     addition at a concentration of 3.
F007 The inoculum source was the Sittingbourne Sewage works in Kent, England,
     and was prepared according to methods described in the OECD 301D
     guideline. The test substance was added to the test medium by direct
     addition at a concentration of 3.0 mg/L. Test systems were incubated at
     20 ± 1 °C and biodegradation was determined by measuring the oxygen
     concentration on days 5, 15, and 28. Each sampling of the test substance
     and control was conducted in duplicate. The theoretical oxygen demand was
     2.82 mg O2 per mg test substance and a theoretical carbon dioxide (CO2)
     evolution of 2.59 mg CO2 per mg test substance. Sodium benzoate was used
     as the positive control.
     The purity of the test substance was not supplied, but the infra-red
     spectrum of the test substance matched a published standard (density =
     0.723 to 0.726 kg/L). The test substance was stored in the dark at
     ambient temperature. Nitrogen was blown over the surface of the material
     when the container was opened and exposed to air in order to minimize
     peroxide formation.
F020 241216
EOR
F002 455
F010 3.5
F004 3
F005 CL
F006 5-day BOD = 0.19 \text{ g/g}, representing 7% biodegradation of the test
     substance.
F007 5-day BOD = 0.19 g/g, representing 7% biodegradation of the test
     substance.
F020 244083
EOR
F002 455
F010 3.5
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* *

Test Substance = 8..85

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F004 3
F005 RE
F006 Bridié A, Wolff C and Winter M (1979). BOD and COD of some
     petrochemicals. Wat. Res. 13, 627-630.
F007 Bridié A, Wolff C and Winter M (1979). BOD and COD of some
    petrochemicals. Wat. Res. 13, 627-630.
F020 244085
EOR
F002 455
F010 3.5
F004 3
F005 RL
F006 The article presented a brief description of the testing methods, but
     cited a reliable guideline method in use at the time of the study.
F007 The article presented a brief description of the testing methods, but
     cited a reliable guideline method in use at the time of the study.
F020 244084
EOR
F002 455
F010 3.5
F004 3
F005 RM
F006 Test type: Biological Oxygen Demand (BOD)
F007 Test type: Biological Oxygen Demand (BOD)
F020 244108
EOR
F002 455
F010 3.5
F004 3
F005 RS
F006 0.19 g 02/g test material at 20 \pm 1°C
     The theoretical oxygen demand (ThOD) of the test substance was 2.82. The
     percent ThOD in 5 days was 7%.
* *
* *
     The article stated that the only deviation from the standard method was
     the addition of 0.5 mg/
F007 0.19 g 02/g test material at 20 \pm 1°C
    The theoretical oxygen demand (ThOD) of the test substance was 2.82. The
    percent ThOD in 5 days was 7%.
* *
    The article stated that the only deviation from the standard method was
     the addition of 0.5 mg/L allylthiourea to prevent nitrification.
F020 244082
EOR
F002 455
F010 3.5
F004 3
F005 TC
F006 The article stated that the test method followed APHA Standard Method
     No. 219 (1971). The test was run at a temperature of 20 \pm 1°C. 500-mL
     test solutions were seeded with a filtered 10-mL volume of the effluent
     from a biological sanitar
F007 The article stated that the test method followed APHA Standard Method
    No. 219 (1971). The test was run at a temperature of 20 \pm 1°C. 500-\text{mL}
     test solutions were seeded with a filtered 10-mL volume of the effluent
     from a biological sanitary waste treatment plant. The activity of the
     inoculum was check by including a treatment containing a mixture of
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glucose and glutamic acid. Test mixtures were stirred using a magnetic
     stirrer.
F020 244081
EOR
F002 455
F010 3.5
F004 4
F005 CL
F006 The authors indicated that K1 values >10 L/g VSS-h represent readily
     biodegradable organic compounds. Based on the results of this study, all
     three test methodologies showed the test substance to be effectively
     utilized by activated sludge
F007 The authors indicated that K1 values >10 L/g VSS-h represent readily
     biodegradable organic compounds. Based on the results of this study, all
     three test methodologies showed the test substance to be effectively
     utilized by activated sludge microorganisms under aerobic conditions.
F020 244090
EOR
F002 455
F010 3.5
F004 4
F005 ME
F006 Comparison study of three aerobic biodegradation methods)
     Continuous Biological Treatment:
     (1) EPA Method 304B (EPA, 1994)
* *
    Batch Methods:
* *
     (2) Batch Oxygen addition (BOX) (Rajagopalan et al., 1998), and
* *
     (3) Serum Bottle Test (SBT) (Rajag
F007 Comparison study of three aerobic biodegradation methods)
* *
     Continuous Biological Treatment:
* *
     (1) EPA Method 304B (EPA, 1994)
* *
     Batch Methods:
* *
     (2) Batch Oxygen addition (BOX) (Rajagopalan et al., 1998), and
     (3) Serum Bottle Test (SBT) (Rajagopalan et al., 1998)
* *
F020 244086
EOR
F002 455
F010 3.5
F004 4
F005 RE
F006 Cano M, Wilcox M and van Compernolle R (1999). A direct comparison of
     U.S. Environmental Protection Agency's Method 304B and batch tests for
     determining activated-sludge biodegradation rate constants for volatile
     organic compounds. Wat. Env
F007 Cano M, Wilcox M and van Compernolle R (1999). A direct comparison of
     U.S. Environmental Protection Agency's Method 304B and batch tests for
     determining activated-sludge biodegradation rate constants for volatile
     organic compounds. Wat. Environ. Res. 71, 1345-1353.
F020 244092
EOR
F002 455
F010 3.5
F004 4
F005 RE
F006 Rajagopalan S, van Compernolle R, Meyer M, Cano M, and Sun P (1998).
     Comparison of methods for determining biodegradation kinetics of volatile
     organic compounds. Water Environ. Res. 70, 291-298.
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F007 Rajagopalan S, van Compernolle R, Meyer M, Cano M, and Sun P (1998).
     Comparison of methods for determining biodegradation kinetics of volatile
     organic compounds. Water Environ. Res. 70, 291-298.
F020 244094
EOR
F002 455
F010 3.5
F004 4
F005 RE
F006 U.S. EPA. (1994). Method 304B - Determination of biodegradation rates of
     organic compounds (scrubber option). Appendix A to Part 63 - Test
     Methods. Fed. Regist. 59,78, 19594.
F007 U.S. EPA. (1994). Method 304B - Determination of biodegradation rates of
     organic compounds (scrubber option). Appendix A to Part 63 - Test
     Methods. Fed. Regist. 59,78, 19594.
F020 244093
EOR
F002 455
F010 3.5
F004 4
F005 RL
F006 The publication presented a well-documented study based on ound
     scientific principles.
F007 The publication presented a well-documented study based on ound
     scientific principles.
F020 244091
EOR
F002 455
F010 3.5
F004 4
F005 RM
F006 Exposure Period:
* *
     Method 304: 30 days
* *
     BOX: 0.5 to 2 hours
* *
     SBT: 0.5 to 2 hours
F007 Exposure Period:
* *
    Method 304: 30 days
* *
     BOX: 0.5 to 2 hours
* *
     SBT: 0.5 to 2 hours
F020 244087
EOR
F002 455
F010 3.5
F004 4
F005 RS
F006 The average percent removal of the test substance in the continuous
     activated sludge unit (EPA Method 304B) was 99.4%.
* *
* *
     Three experimental trial runs with each of the three biodegradation
     methods yielded the following average first-order bi
F007 The average percent removal of the test substance in the continuous
     activated sludge unit (EPA Method 304B) was 99.4%.
* *
     Three experimental trial runs with each of the three biodegradation
     methods yielded the following average first-order biodegradation rate
     constants (K1 = L/g Volatile Suspended Solids-h) for the test substance:
```

```
K1 (L/g VSS-h)
* *
     304B
                 98
* *
     BOX
                 17.4
* *
     SBT
                 19.2
F020 244089
EOR
F002 455
F010 3.5
F004 4
F005 TC
F006 A pilot-scale continuous activated sludge unit served as the source of
    biomass for kinetic rate constant comparisons of the three methods. The
     activated sludge was acclimated in the pilot unit by feeding a synthetic
     cocktail of eight volat
F007 A pilot-scale continuous activated sludge unit served as the source of
    biomass for kinetic rate constant comparisons of the three methods. The
     activated sludge was acclimated in the pilot unit by feeding a synthetic
     cocktail of eight volatile organic compounds during a 2-month
     equilibration period. Equilibration and testing was done at ambient
     temperature (22 to 24°C). The hydraulic retention time (HRT) was 7.7
    hours and the solids retention time (SRT) was 33 days. Average organic
     removal efficiencies based on COD and TOC were 92 and 88%, respectively.
* *
* *
    During the biodegradation testing using Method 304B, feed and effluent
     samples were collected in headspace-free VOA vials and stored at 4°C
     until analyzed. Samples were analyzed by purge-and-trap gas
     chromatography using a flame ionization detector. Triplicate
    biodegradation runs on the test compound were conducted with at least six
     influent and effluent samples taken at 1 HRT (approx. 8 hours) intervals.
* *
    The two batch biodegradation testing methods (BOX and SBT) used activated
     sludge biomass from the pilot-scale reactor. Biomass was diluted using
     effluent from the system to achieve range of 200 to 600 mg/L. The test
     compound was injected into the batch reactors and the concentration was
    monitored over time by collecting gas samples directly from the headspace
     using an automatic sampling pump and analyzing immediately using gas
     chromatography.
F020 244088
EOR
F002 455
F010 3.5
F004 5
F005 CL
F006 Diisopropyl ether can be effectively biodegraded in high biomass aerobic
F007 Diisopropyl ether can be effectively biodegraded in high biomass aerobic
    reactors.
F020 244098
EOR
F002 455
F010 3.5
F004 5
F005 RE
F006 Pruden A, Suidan M, Venosa A and Wilson G (2001). Biodegradaton of methyl
     tert-butyl ether under various substrate conditions. Environ. Sci.
     Technol. 35, 4235-4241.
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F007 Pruden A, Suidan M, Venosa A and Wilson G (2001). Biodegradaton of methyl
     tert-butyl ether under various substrate conditions. Environ. Sci.
     Technol. 35, 4235-4241.
F020 244100
EOR
F002 455
F010 3.5
F004 5
F005 RL
F006 The report provided adequate details of the test conditions but reported
     only a text description of biodegradation results.
F007 The report provided adequate details of the test conditions but reported
     only a text description of biodegradation results.
F020 244099
EOR
F002 455
F010 3.5
F004 5
F005 RM
F006 Inoculum consisted of a mixture of the following:
     1) mixed liquor from the Metropolitan Sewer District (MSD), Cincinnati,
     OH,
* *
     2) mixed liquor from Shell Development Co., Houston, TX, and
     3) aquifer material wash water from a MTBE-contaminat
F007 Inoculum consisted of a mixture of the following:
* *
     1) mixed liquor from the Metropolitan Sewer District (MSD), Cincinnati,
* *
     2) mixed liquor from Shell Development Co., Houston, TX, and
     3) aquifer material wash water from a MTBE-contaminated site in Port
     Hueneme, CA.
F020 244095
EOR
F002 455
F010 3.5
F004 5
F005 RS
F006 The authors indicated that removal of DIPE was comparable to that
     achieved for MTBE, which was greater than 99.9%.
F007 The authors indicated that removal of DIPE was comparable to that
     achieved for MTBE, which was greater than 99.9%.
F020 244097
EOR
F002 455
F010 3.5
F004 5
F005 TC
F006 Bioreactors (Autoclave Engineers, Erie, PA) were initially seeded with 2
     L of mixed liquor from the MSD, 600 mL of mixed liquor from Shell
     Development Co., and 140 mL of aquifer wash water. Cultures were
     maintained on a total influent feed
F007 Bioreactors (Autoclave Engineers, Erie, PA) were initially seeded with 2
     L of mixed liquor from the MSD, 600 mL of mixed liquor from Shell
    Development Co., and 140 mL of aquifer wash water. Cultures were
    maintained on a total influent feed of 417 mg/L chemical oxygen demand
     (COD) divided as 50% methyl tert-butyl ether (MTBE) and 50% as
     diisopropyl ether (DIPE).
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Reactors were well mixed and controlled to a temperature of 20°C. To
     retain high biomass levels, a polyethylene porous pot was inserted into
     the reactor. The pots consisted of 0.45 cm thick filter-grade
     polyethylene (pore size = 20 mm), with an internal diameter of 19.1 cm
     and a height of 29.2 cm. Initially, a solids retention time of 18 days
     was maintained by wasting intentionally from the reactor. Subsequently
     (after about 120 days) intentional wasting ceased and only took place
     during sampling of the reactors.
     The combined influent flow rate was 2.37 L/d, with 80% of the total flow
    provided by a pH-adjustment solution, and 20% provided by an acidified
     nutrient solution. The pH-adjustment solutions contained deionized
     water, MTBE and DIPE fed by a syringe infusion pump, and an appropriate
     amount of 10N sodium hydroxide to maintain the pH between 7.4 and 8.0.
     The nutrient solution consisted of deionized water with essential salts
     and vitamins added to promote biological growth. Final nutrient
     concentrations inside the reactor were as follows: (NH4)2SO4, 93 mg/L;
    MgSO4, 69.6 mg/L; CaCl2o2H2O, 22.5 mg/L; K2HPO4, 6.9 mg/L; CuSO4oH2O,
     0.08 mg/L; Na2MoO4o2H2O, 0.15 mg/L; MnSO4oH2O, 0.13 mg/L; ZnCl2, 0.23
     mg/L; CoCl2o6H2O, 0.42 mg/L; and FeCl2o4H2O, 17.25 mg/L. The hydraulic
    retention time was 4.2 days with a total reactor volume of 9.95 L and an
     enrichment culture volume of 6 L.
* *
    Effluent from the reactors was monitored weekly for the presence of MTBE
    and DIPE using gas chromatography equipped with a flame ionization
     detector (FID) and a 60/80 Carbopack B5% Carbowax 20 M glass column. The
    pH of the system was measured daily, and COD and dissolved organic carbon
     (DOC) was measured weekly.
F020 244096
EOR
F002 455
F010 3.5
F004 6
F005 CL
F006 The test substance was not aerobically biodegraded by indigenous
     subsurface microflora.
F007 The test substance was not aerobically biodegraded by indigenous
     subsurface microflora.
F020 244104
EOR
F002 455
F010 3.5
F004 6
F005 RE
F006 Zenker M, Borden R, and Barlaz M (1999). Investigation of the intrinsic
     biodegradation of alkyl and cyclic ethers. Vol. 1, pp 165-170, 5th Int.
     In Situ On-Site Biorem. Symp.
F007 Zenker M, Borden R, and Barlaz M (1999). Investigation of the intrinsic
    biodegradation of alkyl and cyclic ethers. Vol. 1, pp 165-170, 5th Int.
     In Situ On-Site Biorem. Symp.
F020 244106
EOR
F002 455
F010 3.5
F004 6
F005 RL
F006 The testing method did not follow any specific regulatory guideline
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method, but the publication provided valuable information using sound
     scientific principles.
F007 The testing method did not follow any specific regulatory guideline
     method, but the publication provided valuable information using sound
     scientific principles.
F020 244105
EOR
F002 455
F010 3.5
F004 6
F005 RM
F006 Test type: soil/water microcosm
F007 Test type: soil/water microcosm
F020 244103
EOR
F002 455
F010 3.5
F004 6
F005 RS
F006 No detectable biodegradation of the test substance occurred after one
     year of incubation.
F007 No detectable biodegradation of the test substance occurred after one
    year of incubation.
F020 244102
EOR
F002 455
F010 3.5
F004 6
F005 TC
F006 Soil and water from an aquifer with previous exposure to methyl
     tert-butyl ether (MTBE) was collected using a coring device and a pump.
     The material was brought to the laboratory where the sediment was
     thoroughly mixed. Groundwater was fi
F007 Soil and water from an aquifer with previous exposure to methyl
     tert-butyl ether (MTBE) was collected using a coring device and a pump.
     The material was brought to the laboratory where the sediment was
     thoroughly mixed. Groundwater was filtered through a 0.45 mm filter and
     sparged for 12 hours with sterile air to oxygenate the water and to
    remove background volatile chemicals. Analysis by gas chromatography
     indicated that concentrations of MTBE in the aqueous samples were <10
    mg/L. Microcosms were constructed in amber 255-mL screw-top bottles
     sealed with TeflonÒ MininertÒ valves. Each bottle contained 150 g of wet
     sediment, 140 mL of sterile groundwater and 3000 mg/L of diisopropyl
     ether (DIPE). Treatments were constructed in triplicate with matching
     abiotic controls. Sediment used for the abiotic controls was autoclaved
     for one hour on each of three consecutive days. Additionally, 250 mg/L
     of mercuric chloride was added to ensure no biological activity.
    Microcosms were incubated in the dark at 16 °C.
* *
    All samples were analyzed every 30 days by purge and trap gas
     chromatography and flame ionization detection to determine concentrations
     of the test substance. Test substance disappearance relative to abiotic
     controls was the principal indicator of biodegradation.
F020 244101
EOR
F002 455
```

F010 3.5

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F004 7
F005 CL
F006 Diisopropyl ether was a persistent molecule that resisted anaerobic
     destruction. After 252 days, no evidence for the anaerobic
     biodegradability of diisopropyl ether was obtained.
F007 Diisopropyl ether was a persistent molecule that resisted anaerobic
     destruction. After 252 days, no evidence for the anaerobic
     biodegradability of diisopropyl ether was obtained.
F020 244111
EOR
F002 455
F010 3.5
F004 7
F005 RE
F006 Suflita J and Mormile M (1993). Anaerobic biodegradation of known and
     potential gasoline oxygenates in the terrestrial subsurface. Environ.
     Sci. Technol. 27, 976-978.
F007 Suflita J and Mormile M (1993). Anaerobic biodegradation of known and
    potential gasoline oxygenates in the terrestrial subsurface. Environ.
     Sci. Technol. 27, 976-978.
F020 244113
EOR
F002 455
F010 3.5
F004 7
F005 RL
F006 The publication reported a well-documented study that meets basic
     scientific principles.
F007 The publication reported a well-documented study that meets basic
     scientific principles.
F020 244112
EOR
F002 455
F010 3.5
F004 7
F005 RS
F006 Biodegradation Rate (ppm C/day) = 0
    Methane recovery (% theoretical) = 0
F007 Biodegradation Rate (ppm C/day) = 0
    Methane recovery (% theoretical) = 0
F020 244110
EOR
F002 455
F010 3.5
F004 7
F005 TC
F006 Diisopropyl ether was tested for the ability of the compound to be
     completely biodegraded to methane in an aquifer slurry. Sediment and
     groundwater were collected from a methanogenic portion of a shallow
     anoxic aquifer polluted by municipal
F007 Diisopropyl ether was tested for the ability of the compound to be
     completely biodegraded to methane in an aquifer slurry. Sediment and
     groundwater were collected from a methanogenic portion of a shallow
     anoxic aquifer polluted by municipal landfill leachate. Slurries were
    prepared by placing 50 g of sediment and 75 mL of groundwater in sterile
     160-mL serum bottles. The bottles were sealed with Teflon-lined stoppers
     and incubated in the dark at room temperature. Diisopropyl ether was
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added to the incubation mixture to reach an initial substrate
     concentration of 50 ppm C. Pressure increases resulting from biogas
     formation (CH4 and CO2) were monitored with an automated pressure
     transducer system. The acclimation time was estimated as the amount of
     time where no significant pressure difference was measured between the
     substrate-amended treatment and un-amended controls.
* *
    At the end of the incubation period, biodegradation was measured as the
     depletion of parent substrate and the formation of methane over
    background controls. Measurements were made using gas chromatography
     equipped with a flame ionization detector. A 1.8 m \times 0.32 cm 80/100
    porapak Q column or a 0.2% Carbowax 1500 on Carbopack C column were used
     for headspace methane analyses and test substance determinations,
     respectively. Autoclaved controls were similarly assayed and were
     uniformly unable to exhibit methane formation or test substance
     disappearance. The amount of methane formed in aquifer incubations was
     compared to that theoretically expected based on the Buswell equation.
F020 244109
EOR
F002 455
F010 3.5
F004 8
F005 CL
F006 Diisopropyl ether is not anaerobically degraded under nitrate- or
     sulfate-reducing conditions, and it is not anaerobically degraded under
     methanogenic conditions.
F007 Diisopropyl ether is not anaerobically degraded under nitrate- or
     sulfate-reducing conditions, and it is not anaerobically degraded under
     methanogenic conditions.
F020 244117
EOR
F002 455
F010 3.5
F004 8
F005 RE
F006 Mormile M, Liu S and Suflita J (1994). Anaerobic biodegradation of
     qasoline oxygenates: Extrapolation of information to multiple sites and
     redox conditions. Environ. Sci. Technol. 28, 1727-1732.
F007 Mormile M, Liu S and Suflita J (1994). Anaerobic biodegradation of
     gasoline oxygenates: Extrapolation of information to multiple sites and
     redox conditions. Environ. Sci. Technol. 28, 1727-1732.
F020 244119
E \cap R
F002 455
F010 3.5
F004 8
F005 RL
F006 The publication reported a well-documented study that meets basic
     scientific principles.
F007 The publication reported a well-documented study that meets basic
     scientific principles.
F020 244118
EOR
F002 455
F010 3.5
F004 8
F005 RM
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F006 Exposure period: 85, 180, or 244 days
F007 Exposure period: 85, 180, or 244 days
F020 244114
EOR
F002 455
F010 3.5
F004 8
F005 RS
F006 Biodegradation of diisopropyl ether with sulfate or nitrate available as
     electron acceptors:
                            SO4 or NO3
* *
                       Substrate Amount Consumed
                                                         Rate
* *
                        Loss (%) (% Theoretical) (umol/SO4/
F007 Biodegradation of diisopropyl ether with sulfate or nitrate available as
     electron acceptors:
* *
* *
                            SO4 or NO3
* *
                       Substrate Amount Consumed
                                                         Rate
* *
                        Loss (%) (% Theoretical)
                                                    (umol/SO4/day)
* *
     sulfate-reducing
                           0
                                         0
                                                          0
                                         0
                                                          0
     nitrate-reducing
                           0
* *
* *
* *
     Biodegradation of diisopropyl ether under methanogenic conditions:
* *
                               Degradation
                                              Methane Recovery
* *
                             Rate (ppm C/day)
                                                 (% Expected)
* *
     Fuel-impacted river
* *
                                                        0
       sediment
* *
     Industrial/sewage
* *
       impacted creek sediment
F020 244116
EOR
F002 455
F010 3.5
F004 8
F005 TC
F006 Several tests were carried out to determine the anaerobic biodegradation
     of the test substance. Three experiments were done to determine
     biodegradation under sulfate- and nitrate-reducing conditions and under
     methanogenic conditions. Sedi
F007 Several tests were carried out to determine the anaerobic biodegradation
     of the test substance. Three experiments were done to determine
     biodegradation under sulfate- and nitrate-reducing conditions and under
     methanogenic conditions. Sediment and surface water (or groundwater)
     from three sources were used as inoculum in separate experiments; (1)
     sediment/groundwater from a landfill leachate impacted aquifer, (2)
     sediment/surface water from a river historically impacted by oil storage
     and barge loading facilities, and (3) sediment/surface water from a creek
     impacted by industrial waste and domestic sewage sludge.
* *
     Slurries were prepared by placing 50 g of sediment and 75 mL of water
     into sterile 160-mL serum bottles. Water was amended with sodium sulfide
     (1 mM) and resazurin (0.0002%) to serve as reductant and redox indicator,
     respectively. The bottles were sealed with stoppers and the headspace
     above the slurries was adjusted to 80% N2:20%CO2 (1 atm). To the landfill
     leachate-impacted samples, either sodium sulfate (5mM) or sodium nitrate
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(8 mM) was added in order to assess potential test substance decay
     coupled with the consumption of these electrons (referred to as
     sulfate-reducing and nitrate-reducing incubations, respectively). The
     test substance was added to the slurries to give an initial concentration
     of 50 ppm C. The rates of methane production, sulfate reduction, and
    nitrate depletion were monitored in slurries receiving the test substance
     and compared to test substance-free controls. All incubations were done
     in the dark at 24°C. The sulfate-reducing experiment was run for 244
     days, the nitrate-reducing experiment was run for 85 days, and the
    methanogenic experiment was run for 180 days.
    In the methanogenic incubations, increases in headspace pressure were
    routinely monitored. Parent compound depletion and formation of methane
    were confirmed by gas chromatography (GC). The net amount of sulfate and
    nitrate depletion over the controls was monitored by high pressure liquid
     chromatrography (HPLC).
F020 244115
EOR
F002 455
F010 3.5
F004 9
F005 CL
F006 Diisopropyl ether was degraded by 78% within 24 hours.
F007 Diisopropyl ether was degraded by 78% within 24 hours.
F020 244122
EOR
F002 455
F010 3.5
F004 9
F005 RE
F006 Hernandez-Perez G, Fayolle F and Vandecasteele J-P (2001). Biodegradation
     of ethyl t-butyl ether (ETBE), methyl t-butyl ether (MTBE) and t-amyl
     methyl ether (TAME) by Gordonia terrae. Appl. Microbiol. Biotechnol. 55,
     117-121.
F007 Hernandez-Perez G, Fayolle F and Vandecasteele J-P (2001). Biodegradation
     of ethyl t-butyl ether (ETBE), methyl t-butyl ether (MTBE) and t-amyl
     methyl ether (TAME) by Gordonia terrae. Appl. Microbiol. Biotechnol. 55,
     117-121.
F020 244124
EOR
F002 455
F010 3.5
F004 9
F005 RL
F006 Information on the analytical method was not provided in the report.
F007 Information on the analytical method was not provided in the report.
F020 244123
EOR
F002 455
F010 3.5
F004 9
F005 RS
F006 Diisopropyl ether was degraded by 78% over the 24-hour incubation period.
* *
* *
     Comparison of DIPE biodegradation to ETBE:
* *
        Test Substance Degradation (%)
* *
             ETBE
                               100
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* *
* *
     The authors indicated concentrations of the test substanc
F007 Diisopropyl ether was degraded by 78% over the 24-hour incubation period.
* *
* *
     Comparison of DIPE biodegradation to ETBE:
* *
        Test Substance Degradation (%)
* *
             ETBE
* *
* *
     The authors indicated concentrations of the test substance in the flasks
     were quantified by analytical means. The method was not described in the
     report, but was referenced in an earlier publication by the same workers.
F020 244121
EOR
F002 455
F010 3.5
F004 9
F005 TC
F006 The capacity of ethyl t-butyl ether (ETBE)-induced resting cells of the
     inoculum to degrade diisopropyl ether was tested in sealed flasks. The
     article focused on biodegradation of ETBE, methyl t-butyl ether (MTBE)
     and t-amyl methyl ether (T
F007 The capacity of ethyl t-butyl ether (ETBE)-induced resting cells of the
     inoculum to degrade diisopropyl ether was tested in sealed flasks. The
     article focused on biodegradation of ETBE, methyl t-butyl ether (MTBE)
     and t-amyl methyl ether (TAME), but was tested on other ethers including
    diisopropyl ether (DIPE).
* *
    G. terrae IFP 2001 was cultivated on ETBE-supplemented MM medium. After
     24 hours incubation, bacteria were harvested by centrifugation (20,000 g
     for 20 minutes), washed twice in 100 mM Tris-HCl buffer at pH 7.0 and
     re-suspended in Tris-HCl. The test substance was added to 20-mL cell
     suspensions in 125-mL sealed flasks. Flasks were incubated for 24 hours
     at 30 °C with orbital shaking. Initial cell concentration was 0.5 g/L.
     The test substance was tested at 100 mg/L. Filtered samples were
     analyzed at 0-hour and 24-hours.
F020 244120
EOR
F002 455
F010 3.7
F004 1
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, v3.12. Syracuse
     Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, v3.12. Syracuse
     Research Corporation, Syracuse, NY, USA.
F020 243383
EOR
F002 455
F010 3.7
F004 1
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F007 This robust summary has a reliability rating of 2 because the data are
    calculated and not measured.
F020 243382
EOR
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F002 455
F010 3.7
F004 1
F005 RM
F006 A log bioconcentration factor (BCF) of 0.47 is calculated (BCF = 2.95).
     With respect to a log Kow = 1.52, which was used to calculate the BCF,
     diisopropyl ether in the aquatic environment is expected to have a low
    bioaccumulation potential.
F007 A log bioconcentration factor (BCF) of 0.47 is calculated (BCF = 2.95).
     With respect to a log Kow = 1.52, which was used to calculate the BCF,
     diisopropyl ether in the aquatic environment is expected to have a low
    bioaccumulation potential.
F020 243381
EOR
F002 455
F010 3.7
F004 2
F005 RE
F006 EPI SuiteTM (2000). Estimation Program Interface Suite, v3.12. Syracuse
    Research Corporation, Syracuse, NY, USA.
F007 EPI SuiteTM (2000). Estimation Program Interface Suite, v3.12. Syracuse
    Research Corporation, Syracuse, NY, USA.
F020 243469
EOR
F002 455
F010 3.7
F004 2
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F020 243468
EOR
F002 455
F010 3.7
F004 2
F005 RM
F006 A log bioconcentration factor (BCF) of 1.15 is calculated (BCF = 14.06).
     With respect to a log Kow = 2.4, which was used to calculate the BCF,
     disiopropyl ether in the aquatic environment is expected to have a low
    bioaccumulation potential.
F007 A log bioconcentration factor (BCF) of 1.15 is calculated (BCF = 14.06).
     With respect to a log Kow = 2.4, which was used to calculate the BCF,
     disiopropyl ether in the aquatic environment is expected to have a low
    bioaccumulation potential.
F020 243467
EOR
F002 455
F010 3.8
F004 1
F005 RE
F006 Church C and Tratnyek P (2000). Process level investigations of the in
     situ degradation of MTBE. Presented at the MTBE Biodegradation Workshop,
     February 1-3, 2000. Cincinnati, OH, USA.
F007 Church C and Tratnyek P (2000). Process level investigations of the in
     situ degradation of MTBE. Presented at the MTBE Biodegradation Workshop,
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February 1-3, 2000. Cincinnati, OH, USA.
F020 244127
EOR
F002 455
F010 3.8
F004 1
F005 RE
F006 Eastern Research Group Inc. (2001). Summary of Workshop on Biodegradation
     of MTBE, February 1-3, 2000. Report No. EPA/625/R-01/001A, February,
     2001.
F007 Eastern Research Group Inc. (2001). Summary of Workshop on Biodegradation
     of MTBE, February 1-3, 2000. Report No. EPA/625/R-01/001A, February,
     2001.
F020 244126
EOR
F002 455
F010 3.8
F004 1
F005 RE
F006 Kropp K, Mormile M and Suflita J (2000). Anaerobic biodegradation of MTBE
     and alternative gasoline oxygenates. Presented at the MTBE Biodegradation
     Workshop, February 1-3, 2000. Cincinnati, OH, USA.
F007 Kropp K, Mormile M and Suflita J (2000). Anaerobic biodegradation of MTBE
     and alternative gasoline oxygenates. Presented at the MTBE Biodegradation
     Workshop, February 1-3, 2000. Cincinnati, OH, USA.
F020 244128
EOR
F002 455
F010 3.8
F004 1
F005 RE
F006 Scow K, Smith A, Leung J, Mackay D and Lory E (2000). Bioaugmentation of
     MTBE-contaminated groundwater with bacterial strain PM1. Presented at the
    MTBE Biodegradation Workshop, February 1-3, 2000. Cincinnati, OH, USA.
F007 Scow K, Smith A, Leung J, Mackay D and Lory E (2000). Bioaugmentation of
    MTBE-contaminated groundwater with bacterial strain PM1. Presented at the
     MTBE Biodegradation Workshop, February 1-3, 2000. Cincinnati, OH, USA.
F020 244129
EOR
F002 455
F010 3.8
F004 1
F005 RM
F006 The article reports on a U.S EPA and American Petroleum Institute
     workshop (February 1-3, 2000) on biodegradation of methyl tert-butyl
     ether (MTBE)-contaminated soils and groundwater. MTBE is one of a group
     of structurally similar compound
F007 The article reports on a U.S EPA and American Petroleum Institute
     workshop (February 1-3, 2000) on biodegradation of methyl tert-butyl
     ether (MTBE)-contaminated soils and groundwater. MTBE is one of a group
     of structurally similar compounds commonly called alkyl ether oxygenates
     (AEO) that are added to reformulated gasoline to reduce carbon monoxide
    and ozone emissions. Diisopropyl ether (DIPE) is one type of AEO that is
    used in gasoline along others in this class of chemicals. The workshop
     focused on the status of the current research and understanding on
    biodegradation of MTBE and reported relevant information on the
    biodegradation of DIPE and other AEOs used in reformulated gasoline.
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** Pure microbial cultures have been identified and isolated that have

* demonstrated the capability of utilizing MTBE as a sole carbon and energy

* source under aerobic conditions (Scow et al., 2000). This isolate,

* bacterial strain PM1, was studied by Church and Tratnyek (2000) to

* determine the aerobic degradation pathway of MTBE. In their study, the

* authors confirmed the mineralization of MTBE and determined the

* degradation rates of DIPE and other AEOs were of the same order of

* magnitude as the degradation rates of MTBE (Church and Tratnyek, 2000).

* Their results suggested that similar enzyme systems were responsible for

* all of the reactions.

**

** While the majority of research on anaerobic biodegradation of these

* compounds has been unable to show that MTBE is utilized, a few studies

compounds has been unable to show that MTBE is utilized, a few studies have demonstrated that MTBE and other AEOs may be susceptible to attack under anaerobic conditions. Kropp et al. (2000) studied the anaerobic biodegradation potential of MTBE, DIPE, and other oxygenates in sediment slurries under methanogenic conditions. They found definite evidence in the form of methane and carbon dioxide production to conclude that anaerobic degradation was occurring. The workshop authors concluded that anaerobic biodegradation was a phenomena that was not widespread and extremely difficult for these compounds.

F020 244125 EOR F002 455 F010 3.8 F004 2

F005 RM

F006 Diisopropyl ether (DIPE) is one of a group of similar compounds referred * to as alkyl ether oxygenates (AEO) that are added to reformulated * gasoline. Biodegradation studies with DIPE and other AEO compounds have demonstrated the ability of t

F007 Diisopropyl ether (DIPE) is one of a group of similar compounds referred to as alkyl ether oxygenates (AEO) that are added to reformulated gasoline. Biodegradation studies with DIPE and other AEO compounds have demonstrated the ability of these substances to be consumed by pure strains and mixed cultures of bacteria when incubated under aerobic conditions. Church and Tratnyek (2000) showed that bacterial strain PM1, which had been acclimated to methyl t-butyl ether (MTBE), could mineralize MTBE and demonstrated similar degradation rates for DIPE and other AEOs. They concluded that similar enzymes were responsible for all the degradation reactions. Additional evidence showing the wide spectrum of activity of the bacterial enzyme systems to degrade AEOs was provided by Hernandez-Perez et al. (2001). Using isolated Gordonia terrae (strain IFP 2001) that had been grown on ethyl t-butyl ether (ETBE), degradation of a variety of other AEOs could be achieved. DIPE was degraded 78% within 24 hours in their study (Hernandez-Perez et al. 2001).

* *

** Optimum biodegradation in mixed culture systems occurred when the

* microbial culture is allowed a period of acclimation to the substrate.

* For example, Bridié et al. (1979) measured only 7% consumption of the

* theoretical oxygen demand when DIPE was tested in a 5-day BOD test. In

* contrast, Cano et al. (1999) showed rapid utilization of DIPE when

* activated sludge was conditioned to a cocktail of volatile organic

* compounds for two months. In a continuous flow reactor, DIPE removal

* averaged 99.4%. Cano et al. (1999) also measured high rates of

biodegradation of DIPE when comparing the continuous treatment method

(EPA Method 304B) (EPA, 1994) to two batch treatment methods (BOX and SBT methods; Rajagopalan et al., 1998). Based on the measured rate constants, the authors considered DIPE to be readily biodegradable. Pruden et al. (2001) also observed high removal rates (e.g., 99.9%) of DIPE through a continuous flow reactor system. The performance of the reactors was enhanced when biomass was retained in the reactor, suggesting that a long biomass residence time may be needed for complete mineralization. Biodegradation of DIPE is not always observed in biodegradation assays. Zenker et al. (1999) failed to show biodegradation of DIPE over a 1-year period using indigenous microflora in sediment and water from an aquifer that had been previously exposed to MTBE. Given the apparent need of microbial communities for a period of acclimation to DIPE, it is unlikely that DIPE would be considered readily biodegradable in standard guideline studies. However, the evidence shows that DIPE can be inherently degraded by pure strains of bacteria and mixed enrichments of activated sludge microorganisms. * * While available research shows that DIPE is capable of being biodegraded under aerobic conditions, anaerobic biodegradation is extremely difficult and this substance is considered recalcitrant under those conditions. Suflita et al. (1993) showed no biodegradation of DIPE after 252 days of anaerobic incubation. Substrate was added as 50 ppm C to sediment and groundwater collected from a methanogenic portion of a shallow anoxic aquifer. Similarly, DIPE was evaluated for anaerobic biodegradability under methanogenic conditions as well as sulfate and nitrate-reducing conditions (Mormile et al., 1994). Inocula from three sources (e.g., sediment/groundwater from an aquifer impacted by landfill leachate, sediment/surface water from a river impacted by oil storage, and sediment/surface water from a creek impacted by industrial waste and domestic sewage) were used in separate incubations to assess anaerobic biodegradation in sealed serum bottles. No DIPE biodegradation was measured over incubation periods of 85 days (nitrate-reducing conditions), 180 days (methanogenic conditions), and 244 days (sulfate-reducing conditions). Lack of methane production reported by Suflita and Mormile (1993) does not preclude partial anaerobic biodegradation of DIPE in their studies because only methane was monitored. Kropp et al. (2000) studied the anaerobic biodegradation potential of a number of AEOs including DIPE in sediment slurries under methanogenic conditions. They found evidence in the form of methane and carbon dioxide production to conclude that anaerobic biodegradation was occurring, although the authors stated that anaerobic biodegradation was not a widespread phenomena and extremely difficult for these compounds. F020 244130 EOR F002 455 F010 4.1 F004 1 F005 CL F006 96-hour LC50 = 91.7 mg/L based upon measured values. F007 96-hour LC50 = 91.7 mg/L based upon measured values. F020 241341 EOR F002 455 F010 4.1 F004 1

F005 ME

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F006 The water solubility of the test chemical was obtained from literature or
     determined experimentally. A flow through system using proportional
     diluters and modified continuous mini-diluter system was used for
     maintaining the required test co
F007 The water solubility of the test chemical was obtained from literature or
     determined experimentally. A flow through system using proportional
     diluters and modified continuous mini-diluter system was used for
     maintaining the required test concentrations
* *
    Twenty to twenty-five 30 day-old fish, each weighing approximately 0.12
     g, were randomly divided amongst the test tanks (control and five
     different concentrations) with flow-through dilutor systems.
* *
* *
    Lake Superior water maintained at 25°C ± 1°C was used in the test.
    Routine measures of hardness (EDTA) and total alkalinity of test water
    yielded mean values of 45.5 and 42.2 mg/L as CaCO3, respectively. The
     arithmetic mean of the pH was 7.5 and dissolved oxygen was always greater
     than 60% of saturation.
     Fish were supplied from the United States Environmental Protection
     Agency, Environmental Research Laboratory-Duluth culture. They were not
     fed during the test. Deaths were recorded after 1, 3, 6,12, 24, 48, 72,
     and 96 hours.
F020 241339
EOR
F002 455
F010 4.1
F004 1
F005 RE
F006 Veith G, Call D and Brooke L (1983). Structure-Toxicity Relationships for
     the Fathead Minnow, Pimephales promelas: Narcotic Industrial Chemicals.
     Can. J. Fish. Aquat. Sci. 40, 743-748.
F007 Veith G, Call D and Brooke L (1983). Structure-Toxicity Relationships for
     the Fathead Minnow, Pimephales promelas: Narcotic Industrial Chemicals.
     Can. J. Fish. Aquat. Sci. 40, 743-748.
F008 HEDSET
F009 19-10-1993
F020 235979
EOR
F002 455
F010 4.1
F004 1
F005 RL
F006 This robust summary has a reliability rating of 2 because complete
     information on the analytical results were not available and the study
     was not conducted under GLP.
F007 This robust summary has a reliability rating of 2 because complete
     information on the analytical results were not available and the study
     was not conducted under GLP.
F020 241342
EOR
F002 455
F010 4.1
F004 1
F005 RM
F006 Statistics: Trimmed Spearman-Karber Method
F007 Statistics: Trimmed Spearman-Karber Method
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F020 241343
EOR
F002 455
F010 4.1
F004 1
F005 RM
F006 Test method described in reference.
F007 Test method described in reference.
F008 HEDSET
F009 19-10-1993
F020 235978
EOR
F002 455
F010 4.1
F004 1
F005 RS
F006 96-hour LL50 = 91.7 mg/L based upon measured values
* *
     Analytical method used was GC analysis with Flame Ionization Detection
     (GC-FID), performed on a Hewlett-Packard model 5730A gas chromatograph.
     Concentrations of the test chemical were mea
F007 96-hour LL50 = 91.7 mg/L based upon measured values
* *
* *
    Analytical method used was GC analysis with Flame Ionization Detection
     (GC-FID), performed on a Hewlett-Packard model 5730A gas chromatograph.
     Concentrations of the test chemical were measured daily at each exposure
     level.
F020 241340
EOR
F002 455
F010 4.1
F004 2
F005 CL
F006 The predicted 96 h LC50 value for fish (214.1 mg/L) is in good agreement
     with the experimental 96 h LC50 value for fathead minnow (Pimephales
     promelas) (91.7 mg/L) (Veith et al., Can. J. Fish. Aquat. Sci.,
     40:743-748) and 48 h EC50 value fo
F007 The predicted 96 h LC50 value for fish (214.1 mg/L) is in good agreement
    with the experimental 96 h LC50 value for fathead minnow (Pimephales
    promelas) (91.7 mg/L) (Veith et al., Can. J. Fish. Aquat. Sci.,
     40:743-748) and 48 h EC50 value for Daphnia (190.0 mg/L) (Stephenson
    R.R., Shell Research Limited, Report No. SBGR.83.215).
F020 241204
EOR
F002 455
F010 4.1
F004 2
F006 ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs)
     presented in this program are used to predict the aquatic toxicity of
     chemicals based on their similarity of structure to chemicals for which
     the aquatic toxicity has
F007 ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs)
    presented in this program are used to predict the aquatic toxicity of
     chemicals based on their similarity of structure to chemicals for which
     the aquatic toxicity has been previously measured. Most SAR calculations
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in the ECOSAR Class Program are based upon the octanol/water partition

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coefficient (Kow). SARs have been used by the U.S. Environmental
     Protection Agency since 1981 to predict the aquatic toxicity of new
     industrial chemicals in the absence of test data. SARs are developed for
     chemical classes based on measured test data that have been submitted by
     industry or they are developed by other sources for chemicals with
     similar structures, e.g., phenols. Using the measured aquatic toxicity
    values and estimated Kow values, regression equations can be developed
     for a class of chemicals. Toxicity values for new chemicals may then be
     calculated by inserting the estimated Kow into the regression equation
     and correcting the resultant value for the molecular weight of the
     compound.
    To date, over 150 SARs have been developed for more than 50 chemical
     classes. These chemical classes range from the very large, e.g., neutral
     organics, to the very small, e.g., aromatic diazoniums. Some chemical
     classes have only one SAR, such as acid chlorides, for which only a fish
     96-hour LC50 has been developed. The class with the greatest number of
     SARs is the neutral organics, which has SARs ranging from acute and
     chronic SARs for fish to a 14-day LC50 for earthworms in artificial soil.
     The ECOSAR Class Program is a computerized version of the ECOSAR
    analysis procedures as currently practiced by the Office of Pollution
    Prevention and Toxics (OPPT). It has been developed within the
    regulatory constraints of the Toxic Substances Control Act (TSCA).
     a pragmatic approach to SAR as opposed to a theoretical approach.
F020 241201
EOR
F002 455
F010 4.1
F004 2
F005 RE
F006 ECOSAR v0.99h in EPI Suite (2000). Estimation Program Interface Suite,
     v3.12. Syracuse Research Corporation, Syracuse, NY, USA.
F007 ECOSAR v0.99h in EPI Suite (2000). Estimation Program Interface Suite,
     v3.12. Syracuse Research Corporation, Syracuse, NY, USA.
F020 241206
EOR
F002 455
F010 4.1
F004 2
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F020 241205
EOR
F002 455
F010 4.1
F004 2
F005 RS
F006 LC50, 96 h, for fish = 214.1 \text{ mg/L}
F007 LC50, 96 h, for fish = 214.1 \text{ mg/L}
F020 241203
EOR
F002 455
F010 4.1
F004 2
```

```
F005 TC
F006 Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al,
     1987), log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C
     (SRC PhysProp database) were entered into the program.
     Class: Neutral organics
F007 Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al,
     1987), log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C
     (SRC PhysProp database) were entered into the program.
* *
    Class: Neutral organics
F020 241202
EOR
F002 455
F010 4.1
F004 2
F005 TS
F006 Diisopropyl Ether (CAS No. 108-20-3)
F007 Diisopropyl Ether (CAS No. 108-20-3)
F020 241214
EOR
F002 455
F010 4.1
F004 3
F005 CL
F006 96-hour LC50 = 786 \text{ mg/L} based on mean measured values.
     96-hour EC50 = 476 mg/L based on mean measured values.
F007 96-hour LC50 = 786 \text{ mg/L} based on mean measured values.
     96-hour EC50 = 476 mg/L based on mean measured values.
F020 243473
EOR
F002 455
F010 4.1
F004 3
F005 ME
F006 Test solutions were prepared using a proportional diluter system without
    replication. This system provided control and five test substance
     concentrations to glass test vessels. Each vessel held 2 L of test
     solution and the diluter flow ra
F007 Test solutions were prepared using a proportional diluter system without
    replication. This system provided control and five test substance
     concentrations to glass test vessels. Each vessel held 2 L of test
     solution and the diluter flow rate was sufficient to provide 18 volume
    additions per day. An aqueous stock solution of 1050 mg/L was used by
     the diluter to prepare the exposure series. Dilution water was filtered
     Lake Superior water. Typical ranges of water quality factors measured in
     this water were pH (7.4 - 8.2), total hardness (44 - 53 \text{ mg/L as CaCO3}),
     and specific conductance (78 - 86 mmhos/cm).
* *
    Test fish originated from in-house cultures of P. promelas at the U.S.
    EPA Environmental Research Laboratory - Duluth. Fish were not fed 24 h
    prior to testing or during the test. At test initiation, fish were
    randomly placed in test vessels until each vessel contained 10
     individuals. Individuals used in testing were 34 d old and measured 19.0
    mm mean length (SD = 2.534) and 0.104 g mean weight (SD = 0.0433).
    Biomass loading was 1.04 g/L. Death was the major test endpoint.
    Numbers of dead fish were counted daily and any dead fish were removed
     from the vessels. Abnormal behavioral changes were recorded at each
     observation time. LC50 (lethality) and EC50 (total effect) values were
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determined.
* *
     Temperature, dissolved oxygen, and pH were measured daily in all test
     chambers. Mean values (and Standard Deviation) were 24.9 °C (SD = 0.52),
     7.3 mg/L (SD = 0.13), and 7.75 (SD = 0.16), respectively. Total hardness
     and alkalinity were measured once in the control, low, medium, and high
     test levels. Mean values were 43.7 mg/L total hardness as CaCO3 (SD =
     0.96) and 49.6 mg/L alkalinity as CaCO3 (SD = 0.25). Lighting was
    provided by fluorescent bulbs that produced 28 to 48 lumens/sq ft at the
     water surface. The photoperiod was 16 h light and 8 h dark.
    Test substance concentrations were verified in most cases daily during
     the test using gas-liquid chromatography. Concentrations were averaged
     and a mean percent recovery was calculated. The nominal with measured
     concentrations in parentheses were, control (not detected), 157 mg/L (131
     mg/L), 242 mg/L (210 mg/L), 373 mg/L (382 mg/L), 574 mg/L (594 mg/L), and
     883 mg/L (1044 mg/L). The overall percent recovery was 98.7%.
F020 243470
EOR
F002 455
F010 4.1
F004 3
F005 RE
F006 Geiger D, Poirier S, Brooke L and Call D (eds.) (1986). Acute Toxicities
     of Organic Chemicals to Fathead Minnows (Pimephales promelas), Vol. 3.
     Center for Lake Superior Environmental Studies, Univ. of Wisconsin-
     Superior, Superior, WI, USA.
F007 Geiger D, Poirier S, Brooke L and Call D (eds.) (1986). Acute Toxicities
     of Organic Chemicals to Fathead Minnows (Pimephales promelas), Vol. 3.
     Center for Lake Superior Environmental Studies, Univ. of Wisconsin-
     Superior, Superior, WI, USA.
F020 243474
EOR
F002 455
F010 4.1
F004 3
F005 RM
F006 Statistics: LC/EC50 values determined by Trimmed Spearman-Karber Method
F007 Statistics: LC/EC50 values determined by Trimmed Spearman-Karber Method
F020 243471
EOR
F002 455
F010 4.1
F004 3
F005 RS
F006 96-hour LC50 = 786 mg/L based on mean measured values.
     96-hour EC50 = 476 mg/L based on mean measured values.
* *
* *
     The EC50 value was based on mortality and the following abnormal effects:
     loss of schooling behavior, swimming near the surface,
F007 96-hour LC50 = 786 mg/L based on mean measured values.
     96-hour EC50 = 476 mg/L based on mean measured values.
* *
     The EC50 value was based on mortality and the following abnormal effects:
     loss of schooling behavior, swimming near the surface, hypoactive,
     under-reactive to external stimuli, loss of equilibrium.
F020 243472
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EOR
F002 455
F010 4.1
F004 4
F005 CL
F006 96-h LC50 = 900 mg/L based on measured concentrations
F007 96-h LC50 = 900 mg/L based on measured concentrations
F020 243478
EOR
F002 455
F010 4.1
F004 4
F005 ME
F006 Test solutions were prepared using a continuous-flow diluter delivery
     system, which delivered four test substance concentrations and control
     solutions to duplicate test vessels. Dilution water was filtered Lake
     Superior water. Average val
F007 Test solutions were prepared using a continuous-flow diluter delivery
     system, which delivered four test substance concentrations and control
     solutions to duplicate test vessels. Dilution water was filtered Lake
     Superior water. Average values for water quality factors for the
     dilution water were: hardness (44.6 mg/L as CaCO3), total alkalinity
     (44.0 mg/L as CaCO3), and pH (7.6). Test chambers were glass vessels and
     contained 2 L of test solution. Solution flow rates through the test
     chambers was sufficient to provided at least a 95% replacement in
     approximately 4 h. Test substance concentrations were verified daily
     during the test using either gas chromatography or high pressure liquid
     chromatography methods.
    The mean temperature for the test was 25 \pm 0.5°C, and dissolved oxygen
     remained at or above 80% saturation. Lighting was provided by wide
     spectrum fluorescent bulbs at an intensity of 22 to 38 lumens/sq ft over
     the test chambers. The photoperiod was 16 h light and 8 h dark with a
     30-min dusk/dawn transition period.
* *
    Test fish originated from cultures maintained by the U.S. EPA
     Environmental Research Laboratory - Duluth, MN. and were 28 to 34 days
     old (weighing approximately 0.12 g) at the time of testing. A total of
     20 fish per treatment (10/replicate) was used in the test. Fish were
     added to the test chambers 2-3 h before introduction of the test
     solutions. Fish were not fed 24 h before or during the test.
    Mortalities were recorded daily.
F020 243475
EOR
F002 455
F010 4.1
F004 4
F006 Broderius S, and Kahl M (1985). Acute toxicity of organic chemical
     mixtures to the fathead minnow. Aquat. Toxicol. 6:307-322.
F007 Broderius S, and Kahl M (1985). Acute toxicity of organic chemical
     mixtures to the fathead minnow. Aquat. Toxicol. 6:307-322.
F020 243479
EOR
F002 455
F010 4.1
F004 4
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F005 RM
F006 Statistics: Trimmed Spearman-Karber Method or log-probit method.
F007 Statistics: Trimmed Spearman-Karber Method or log-probit method.
F020 243476
EOR
F002 455
F010 4.1
F004 4
F005 RS
F006 96-h LC50 = 900 mg/L based on measured concentrations
     95\% CL = 881 - 920 mg/L
F007 96-h LC50 = 900 mg/L based on measured concentrations
     95\% CL = 881 - 920 mg/L
F020 243477
EOR
F002 455
F010 4.1
F004 5
F005 CL
F006 24-hour LC50 = 380 \text{ mg/L}.
F007 24-hour LC50 = 380 \text{ mg/L}.
F020 243483
EOR
F002 455
F010 4.1
F004 5
F005 ME
F006 The test consisted of exposing groups of six fish to a series of
     concentrations of the test substance for 24 h. Fish were exposed in an
     all glass tanks holding 25 liters of test solution. Dilution water was
     local tap water having the foll
F007 The test consisted of exposing groups of six fish to a series of
     concentrations of the test substance for 24 h. Fish were exposed in an
     all glass tanks holding 25 liters of test solution. Dilution water was
     local tap water having the following characteristics (all values in mg/L):
     C1^- = 65; N02^- = 0; N03^- = 4; S04^-2 = 35; P04^-3 = 0.15; HC03^- = 25; Si02
     = 25; NH4+ = 0; Fe = 0.05; Mn = 0; Ca+2 = 100; Mq+2 = 8; alkali as Na+ = 100
     30; pH = 7.8.
* *
     The test was run at a temperature of 20±1°C, and the solutions were not
     aerated during the test period.
* *
     Test fish had a mean length of 6.2 \pm 0.7 cm, a mean weight of 3.3 \pm 1.0 g
     and were in good health at the time of testing.
* *
     Exposure concentrations were confirmed either by total organic carbon
     analysis or by extraction and subsequent analysis by gas chromatography.
     Measured concentrations were not reported in this study.
F020 243480
EOR
F002 455
F010 4.1
F004 5
F005 RE
F006 Bridie A, Wolff C and Winter M (1979). The acute toxicity of some
     petrochemicals to goldfish. Water Res. 13:623-626.
F007 Bridie A, Wolff C and Winter M (1979). The acute toxicity of some
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petrochemicals to goldfish. Water Res. 13:623-626.
F020 243485
EOR
F002 455
F010 4.1
F004 5
F005 RT
F006 The test was run for only 24 hours to ensure that the dissolved oxygen
     content did not fall below 4 mg/L. The report lacked sufficient detail
     for assessment. It was not stated whether results were based on nominal
     or measured values.
F007 The test was run for only 24 hours to ensure that the dissolved oxygen
     content did not fall below 4 mg/L. The report lacked sufficient detail
     for assessment. It was not stated whether results were based on nominal
     or measured values.
F020 243484
EOR
F002 455
F010 4.1
F004 5
F005 RM
F006 Determination of LC50 by graphical interpolation of log concentrations
     versus percent mortality (APHA, 1971).
F007 Determination of LC50 by graphical interpolation of log concentrations
     versus percent mortality (APHA, 1971).
F020 243481
EOR
F002 455
F010 4.1
F004 5
F005 RS
F006\ 24-hour\ LC50 = 380\ mg/L
* *
* *
     The analytical method was either total organic carbon analysis or gas
     chromatography. It was not reported what method was employed for this
     test substance nor if the result was based on measured concentrations.
F007 24-hour LC50 = 380 mg/L
* *
     The analytical method was either total organic carbon analysis or gas
     chromatography. It was not reported what method was employed for this
     test substance nor if the result was based on measured concentrations.
F020 243482
E \cap R
F002 455
F010 4.1
F004 6
F005 CL
F006 96-hour LC50 = 7,000 mg/L based on nominal concentrations
F007 96-hour LC50 = 7,000 mg/L based on nominal concentrations
F020 243489
EOR
F002 455
F010 4.1
F004 6
F005 ME
F006 The test consisted of exposing groups of fish to a four-dilution series
     of the test substance for a period of 96 h. Test vessels were all glass
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5-gallon aquaria. The volume of test solution was adjusted to assure
     that a biomass loading wa
F007 The test consisted of exposing groups of fish to a four-dilution series
     of the test substance for a period of 96 h. Test vessels were all glass
     5-gallon aquaria. The volume of test solution was adjusted to assure
     that a biomass loading was no more than 1 q fish /liter solution.
     Dilution water was well water having a typical pH of 7.6 to 7.9 and a
    hardness of 55 mg/L (as CaCO3).
* *
     Fish were obtained from a commercial source and assessed for health
     during a 14-d acclimation period prior to testing. During that time they
     were maintained on a commercial fish food diet supplemented with minced
     frozen shrimp. Fish were not fed 48 hours prior to testing. Fish were
     randomly selected for testing and were approximately 33 to 75 mm in
     length.
     The test was run at 23°C. Test solutions were not aerated for the
     initial 24 h, but aeration was applied thereafter if the dissolved oxygen
     concentration was being depleted. Dissolved oxygen readings were taken
     daily, and pH was measured at the end of the test. However, these data
    were not provided in the report.
* *
    Mortality was assessed daily and any dead fish were removed at each
     observation time.
F020 243486
EOR
F002 455
F010 4.1
F004 6
F005 RE
F006 Dawson G, Jennings A, Drozdowski D and Rider E (1975/77). The acute
     toxicity of 47 industrial chemicals to fresh and saltwater fishes. J.
     Haz. Mat. 1:303-318.
F007 Dawson G, Jennings A, Drozdowski D and Rider E (1975/77). The acute
     toxicity of 47 industrial chemicals to fresh and saltwater fishes. J.
    Haz. Mat. 1:303-318.
F020 243491
EOR
F002 455
F010 4.1
F004 6
F005 RI
F006 Documentation was insufficient for evaluation. Basic water quality data
     during the test were not provided. The authors stated that aeration of
     the test solutions was used after 24 hours to ensure maintenance of
     dissolved oxygen. No analyt
F007 Documentation was insufficient for evaluation. Basic water quality data
     during the test were not provided. The authors stated that aeration of
     the test solutions was used after 24 hours to ensure maintenance of
     dissolved oxygen. No analytical verification of exposure concentrations
     were made.
F020 243490
EOR
F002 455
F010 4.1
F004 6
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F005 RM

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F006 The LC50 was determined by plotting survival percentages on
     semi-logarithmic paper and drawing a straight line fit through or near
     significant points above and below 50% survival.
F007 The LC50 was determined by plotting survival percentages on
     semi-logarithmic paper and drawing a straight line fit through or near
     significant points above and below 50% survival.
F020 243487
EOR
F002 455
F010 4.1
F004 6
F005 RS
F006 96-hour LC50 = 7,000 mg/L
     The mortality pattern reported for the test substance suggests that a
     more likely estimate of the LC50 value would lie between 7,900 and 10,000
     mg/L, rather than 7,000 mg/L. This was due to a non-monotonic dose re
F007 96-hour LC50 = 7,000 mg/L
* *
     The mortality pattern reported for the test substance suggests that a
* *
    more likely estimate of the LC50 value would lie between 7,900 and 10,000
    mg/L, rather than 7,000 mg/L. This was due to a non-monotonic dose
    response pattern of mortality. The report authors indicated that the LC50
     value was higher than the published solubility for the test substance.
F020 243488
EOR
F002 455
F010 4.1
F004 7
F005 CL
F006 96-hour LC50 = 6600 mg/L based on nominal concentrations.
F007 96-hour LC50 = 6600 mg/L based on nominal concentrations.
F020 243495
EOR
F002 455
F010 4.1
F004 7
F005 ME
F006 The test consisted of exposing groups of fish to a four-dilution series
     of the test substance for a period of 96 h. Test vessels were all glass
     5-gallon aquaria. The volume of test solution was adjusted to assure
     that a biomass loading wa
F007 The test consisted of exposing groups of fish to a four-dilution series
     of the test substance for a period of 96 h. Test vessels were all glass
     5-gallon aquaria. The volume of test solution was adjusted to assure
     that a biomass loading was no more than 1 g fish /liter solution.
     Dilution water was prepared by adding "instant ocean" salts to well water
     (pH of 7.6 to 7.9; hardness 55 mg/L (as CaCO3)) until a specific gravity
     of 1.018 was achieved.
     Fish were field collected in nets from Horseshoe Bay at Sandy Hook, New
     Jersey. They were held for a 14-d acclimation period prior to testing and
    assessed for health during that time. During the acclimation period they
    were fed minced frozen shrimp. Fish were not fed 48 hours prior to
     testing. Fish were randomly selected for testing and were approximately
     40 to 100 mm in length.
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The test was run at 20°C, and test solutions were continuously aerated
     during the exposure period. Dissolved oxygen readings were taken daily,
     and pH was measured at the end of the test. However, these data were not
    provided in the report.
* *
    Mortality was assessed daily and any dead fish were removed at each
     observation time.
F020 243492
EOR
F002 455
F010 4.1
F004 7
F005 RE
F006 Dawson G, Jennings A, Drozdowski D and Rider E (1975/77). The acute
     toxicity of 47 industrial chemicals to fresh and saltwater fishes. J.
     Haz. Mat. 1:303-318.
F007 Dawson G, Jennings A, Drozdowski D and Rider E (1975/77). The acute
     toxicity of 47 industrial chemicals to fresh and saltwater fishes. J.
     Haz. Mat. 1:303-318.
F020 243497
EOR
F002 455
F010 4.1
F004 7
F005 RT
F006 Documentation was insufficient for evaluation. Basic water quality data
     during the test were not provided. The authors stated that aeration of
     the test solutions was used after 24 hours to ensure maintenance of
     dissolved oxygen. No analyt
F007 Documentation was insufficient for evaluation. Basic water quality data
     during the test were not provided. The authors stated that aeration of
     the test solutions was used after 24 hours to ensure maintenance of
     dissolved oxygen. No analytical verification of exposure concentrations
     were made.
F020 243496
EOR
F002 455
F010 4.1
F004 7
F005 RM
F006 LC50 determined by graphical interpolation of the logarithm of the
     concentration versus the percentage mortality.
F007 LC50 determined by graphical interpolation of the logarithm of the
     concentration versus the percentage mortality.
F020 243493
EOR
F002 455
F010 4.1
F004 7
F005 RS
F006 96-hour LC50 = 6600 mg/L
     The mortality pattern reported for the test substance does not correspond
    with the estimated LC50 value. Given the dose-response pattern, the LC50
     value would lie between 3,200 and 5,000 mg/L, rather than 6,600 mg
F007 96-hour LC50 = 6600 mg/L
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The mortality pattern reported for the test substance does not correspond
    with the estimated LC50 value. Given the dose-response pattern, the LC50
     value would lie between 3,200 and 5,000 mg/L, rather than 6,600 mg/L.
     The authors reported that the result was higher than the reported water
     solubility of the test substance.
F020 243494
EOR
F002 455
F010 4.2
F004 1
F005 CL
F006 After Daphnia magna were exposed to test solutions of di-isopropyl ether
     for 48 hours in a static test, the 24 h and 48 h EC50 values were
     calculated to be 240 mg/L and 190 mg/L, respectively.
F007 After Daphnia magna were exposed to test solutions of di-isopropyl ether
     for 48 hours in a static test, the 24 h and 48 h EC50 values were
     calculated to be 240 mg/L and 190 mg/L, respectively.
F020 241346
EOR
F002 455
F010 4.2
F004 1
F005 RE
F006 Stephenson R (1983). Isopropyl Ether: Acute Toxicity to Daphnia magna and
     Selenastrum capricornutum. Group Research Limited, Report No.
     SBGR.83.215. Shell Research Ltd. Sittingbourne Research Centre,
     Sittingbourne, UK.
F007 Stephenson R (1983). Isopropyl Ether: Acute Toxicity to Daphnia magna and
     Selenastrum capricornutum. Group Research Limited, Report No.
     SBGR.83.215. Shell Research Ltd. Sittingbourne Research Centre,
     Sittingbourne, UK.
F008 HEDSET
F009 19-10-1993
F020 235981
EOR
F002 455
F010 4.2
F004 1
F005 RL
F006 This robust summary has a reliability rating of 2 because it did not
     analytically verify exposure concentrations and the results are based on
    nominal values.
F007 This robust summary has a reliability rating of 2 because it did not
     analytically verify exposure concentrations and the results are based on
     nominal values.
F020 241348
EOR
F002 455
F010 4.2
F004 1
F005 RM
F006 Statistics:
     Probit analysis after log transformation of the concentrations (Finney,
* *
     Probit Analysis, Finney, D.J., Cambridge University Press, 3rd edition,
     p333 (1971)
F007 Statistics:
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1971)
* *
     Probit Analysis, Finney, D.J., Cambridge University Press, 3rd edition,
     p333 (1971)
F020 241349
EOR
F002 455
F010 4.2
F004 1
F005 RS
F006 The 24 h and 48 h Effect Concentration (EC50) values were calculated to
     be 240 mg/L (95% fiducial limits 210 to 280 mg/L) and 190 mg/L (95%
     fiducial limits 160 to 220 mg/L), respectively.
     The immobilization (%) of Daphnia magna (n=10/replic
* *
F007 The 24 h and 48 h Effect Concentration (EC50) values were calculated to
     be 240 mg/L (95% fiducial limits 210 to 280 mg/L) and 190 mg/L (95%
     fiducial limits 160 to 220 mg/L), respectively.
* *
     The immobilization (%) of Daphnia magna (n=10/replicate) are as follows:
* *
* *
     Test Substance
                          Immobilization (%)*
* *
      Loading Rate
* *
                            24 hr
                                       48 hr
            (mg/L)
* *
* *
                             0
                                        0
        0 (control)
* *
                             0
                                        0
             46
* *
             99
                             3
                                        7
* *
            210
                            27
                                       57
* *
            460
                           100
                                      100
           1000
                            100
                                      100
* *
     *mean of 3 replicates
F020 241345
EOR
F002 455
F010 4.2
F004 1
F005 TC
F006 A 48 hour static toxicity test was carried out without renewal of the
     test solutions. Quantities of stock solutions of di-isopropyl ether in
     acetone were added in triplicate sets of 110 mL glass flasks so that when
     made up with reconstitute
F007 A 48 hour static toxicity test was carried out without renewal of the
     test solutions. Quantities of stock solutions of di-isopropyl ether in
     acetone were added in triplicate sets of 110 mL glass flasks so that when
     made up with reconstituted freshwater, an approximately logarithmic
     series of concentrations ranging from 46 to 1000 mg/L was produced. Three
     flasks served as controls and received no test substance. The
     concentration of acetone in all control and test flasks was 0.1 mL/L.
     Precautions were taken to (a) minimise evaporative loss of the test
     substance by use of glass cover slips over the vessel necks and (b) to
     minimize the risk of organisms becoming trapped at the surface by placing
     black paper caps over the flasks to create a darkened zone which the
     organisms would avoid.
     The test temperatures were in the range 20 ± 2°C, pH was in the range 8.2
     to 8.4, the total hardness was 164 mg/L as CaCO3, and dissolved oxygen
     was in the range 8.2 to 9.2 mg/L.
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The daphnids were cultured in-house, derived from a strain obtained (via ICI Brixham Laboratory) from Institut National de Recherche Chimique

Probit analysis after log transformation of the concentrations (Finney,

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Applique (I.R.Ch.A.), France. Age was <24 hours old from 15 to 35 day old
     parents.
     All concentrations of test substance are expressed in terms of quantities
     initially added to the test vessels.
F020 241344
EOR
F002 455
F010 4.2
F004 1
F005 TS
F006 Diisopropyl Ether (CAS No. 108-20-3)
F007 Diisopropyl Ether (CAS No. 108-20-3)
F020 241347
EOR
F002 455
F010 4.2
F004 2
F005 CL
F006 The predicted 48 h LC50 value for Daphnia (221.9 mg/L) is in close
     agreement with the experimental 48 h EC50 value for Daphnia (190.0 mg/L)
     (Stephenson R.R., Shell Research Limited, Report No. SBGR.83.215).
F007 The predicted 48 h LC50 value for Daphnia (221.9 mg/L) is in close
     agreement with the experimental 48 h EC50 value for Daphnia (190.0 mg/L)
     (Stephenson R.R., Shell Research Limited, Report No. SBGR.83.215).
F020 241211
EOR
F002 455
F010 4.2
F004 2
F005 ME
F006 ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs)
     presented in this program are used to predict the aquatic toxicity of
     chemicals based on their similarity of structure to chemicals for which
     the aquatic toxicity has
F007 ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs)
     presented in this program are used to predict the aquatic toxicity of
     chemicals based on their similarity of structure to chemicals for which
     the aquatic toxicity has been previously measured. Most SAR calculations
     in the ECOSAR Class Program are based upon the octanol/water partition
     coefficient (Kow). SARs have been used by the U.S. Environmental
     Protection Agency since 1981 to predict the aquatic toxicity of new
     industrial chemicals in the absence of test data. SARs are developed for
     chemical classes based on measured test data that have been submitted by
     industry or they are developed by other sources for chemicals with
     similar structures, e.g., phenols. Using the measured aquatic toxicity
     values and estimated Kow values, regression equations can be developed
     for a class of chemicals. Toxicity values for new chemicals may then be
     calculated by inserting the estimated Kow into the regression equation
     and correcting the resultant value for the molecular weight of the
     compound.
* *
     To date, over 150 SARs have been developed for more than 50 chemical
     classes. These chemical classes range from the very large, e.g., neutral
     organics, to the very small, e.g., aromatic diazoniums. Some chemical
     classes have only one SAR, such as acid chlorides, for which only a fish
     96-hour LC50 has been developed. The class with the greatest number of
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SARs is the neutral organics, which has SARs ranging from acute and

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chronic SARs for fish to a 14-day LC50 for earthworms in artificial soil.
     The ECOSAR Class Program is a computerized version of the ECOSAR
     analysis procedures as currently practiced by the Office of Pollution
     Prevention and Toxics (OPPT). It has been developed within the
    regulatory constraints of the Toxic Substances Control Act (TSCA).
     a pragmatic approach to SAR as opposed to a theoretical approach.
F020 241207
EOR
F002 455
F010 4.2
F004 2
F005 RE
F006 ECOSAR v0.99h in EPI Suite (2000). Estimation Program Interface Suite,
    v3.12. Syracuse Research Corporation, Syracuse, NY, USA.
F007 ECOSAR v0.99h in EPI Suite (2000). Estimation Program Interface Suite,
     v3.12. Syracuse Research Corporation, Syracuse, NY, USA.
F020 241213
EOR
F002 455
F010 4.2
F004 2
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F020 241212
EOR
F002 455
F010 4.2
F004 2
F005 RS
F006 EC50, 48 h, for Daphnia = 221.9 mg/L
F007 EC50, 48 h, for Daphnia = 221.9 mg/L
F020 241209
EOR
F002 455
F010 4.2
F004 2
F005 TC
F006 Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al,
     1987), log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C
     (SRC PhysProp database) were entered into the program.
    Class: Neutral organics
F007 Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al,
     1987), log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C
     (SRC PhysProp database) were entered into the program.
* *
    Class: Neutral organics
F020 241208
EOR
F002 455
F010 4.2
F004 2
F005 TS
F006 Diisopropyl Ether, CAS No. 108-20-3
F007 Diisopropyl Ether, CAS No. 108-20-3
F020 241210
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EOR
F002 455
F010 4.3
F004 2
F005 CL
F006 The predicted 96 h EC50 value for algae (134.9 mg/L) is in the same range
     as the predicted 48 h LC50 value for Daphnia (221.9 mg/L) and the
    predicted 96 h LC50 value for fish (214.1 mg/L). There is also good
     comparison between the predicted
F007 The predicted 96 h EC50 value for algae (134.9 mg/L) is in the same range
     as the predicted 48 h LC50 value for Daphnia (221.9 mg/L) and the
    predicted 96 h LC50 value for fish (214.1 mg/L). There is also good
     comparison between the predicted and experimental EC50 values for Daphnia
     (221.9 mg/l v 190.0 mg/L, respectively) and for fish (214.1 mg/l v 91.7
     mg/L, respectively).
F020 241198
EOR
F002 455
F010 4.3
F004 2
F005 ME
F006 ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs)
     presented in this program are used to predict the aquatic toxicity of
     chemicals based on their similarity of structure to chemicals for which
     the aquatic toxicity has
F007 ECOSAR version 0.99h, US EPA. The structure-activity relationships (SARs)
    presented in this program are used to predict the aquatic toxicity of
     chemicals based on their similarity of structure to chemicals for which
     the aquatic toxicity has been previously measured. Most SAR calculations
     in the ECOSAR Class Program are based upon the octanol/water partition
     coefficient (Kow). SARs have been used by the U.S. Environmental
     Protection Agency since 1981 to predict the aquatic toxicity of new
     industrial chemicals in the absence of test data. SARs are developed for
     chemical classes based on measured test data that have been submitted by
     industry or they are developed by other sources for chemicals with
     similar structures, e.g., phenols. Using the measured aquatic toxicity
     values and estimated Kow values, regression equations can be developed
     for a class of chemicals. Toxicity values for new chemicals may then be
     calculated by inserting the estimated Kow into the regression equation
     and correcting the resultant value for the molecular weight of the
     compound.
     To date, over 150 SARs have been developed for more than 50 chemical
     classes. These chemical classes range from the very large, e.g., neutral
     organics, to the very small, e.g., aromatic diazoniums. Some chemical
     classes have only one SAR, such as acid chlorides, for which only a fish
     96-hour LC50 has been developed. The class with the greatest number of
     SARs is the neutral organics, which has SARs ranging from acute and
     chronic SARs for fish to a 14-day LC50 for earthworms in artificial soil.
     The ECOSAR Class Program is a computerized version of the ECOSAR
     analysis procedures as currently practiced by the Office of Pollution
     Prevention and Toxics (OPPT). It has been developed within the
     regulatory constraints of the Toxic Substances Control Act (TSCA).
     a pragmatic approach to SAR as opposed to a theoretical approach.
F020 241195
EOR
F002 455
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F010 4.3
F004 2
F005 RE
F006 ECOSAR v0.99h in EPI Suite (2000). Estimation Program Interface Suite,
     v3.12. Syracuse Research Corporation, Syracuse, NY, USA.
F007 ECOSAR v0.99h in EPI Suite (2000). Estimation Program Interface Suite,
     v3.12.
             Syracuse Research Corporation, Syracuse, NY, USA.
F020 241200
EOR
F002 455
F010 4.3
F004 2
F005 RL
F006 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F007 This robust summary has a reliability rating of 2 because the data are
     calculated and not measured.
F020 241199
EOR
F002 455
F010 4.3
F004 2
F005 RS
F006 EC50, 96 h, for green algae = 134.9 \text{ mg/L}
* *
     ChV, 96 h, for green algae = 10.2 mg/L
F007 EC50, 96 h, for green algae = 134.9 \text{ mg/L}
**
     ChV, 96 h, for green algae = 10.2 mg/L
F020 241197
EOR
F002 455
F010 4.3
F004 2
F005 TC
F006 Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al,
     1987), log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C
     (SRC PhysProp database) were entered into the program.
     Class: Neutral organics
F007 Experimental water solubility, 8800 mg/l @ 20°C (Heitmann, W et al,
     1987), log Kow, 1.52 (Funasaki, N et al, 1985) and melting point, -86.6°C
     (SRC PhysProp database) were entered into the program.
* *
     Class: Neutral organics
F020 241196
EOR
F002 455
F010 4.3
F004 2
F005 TS
F006 Diisopropyl Ether (CAS No. 108-20-3)
F007 Diisopropyl Ether (CAS No. 108-20-3)
F020 241215
EOR
F002 455
F010 4.3
F004 3
F005 CL
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F006 96-hour EC50 = >1000 mg/L based on nominal concentrations.
F007 96-hour EC50 = >1000 mg/L based on nominal concentrations.
F020 243500
EOR
F002 455
F010 4.3
F004 3
F005 ME
F006 A 4 d algal growth study was carried out using 10 concentrations of the
     test substance and a control. The test design included six control
     replicates and single vessels dosed with different concentrations of the
     test substance. 250-mL gla
F007 A 4 d algal growth study was carried out using 10 concentrations of the
     test substance and a control. The test design included six control
    replicates and single vessels dosed with different concentrations of the
     test substance. 250-mL glass Erlenmeyer flasks served as the test
     vessels and held 50 mL of culture medium. Culture medium was prepare
     following the recipe given by Miller and Green (1978) with the following
     exceptions; 1) boric acid concentration = 105 mg/L, and 2) sodium
    bicarbonate concentration = 50 mg/L.
* *
    To 10 flasks, quantities of a test substance stock solution made up in
     acetone were added to give a logarithmic series of concentrations ranging
     from 1 to 1000 mg/L (1.0, 2.2, 4.6, 10, 22, 46, 100, 220, 460, and 1000
    mg/L). The concentration of acetone in all flasks including controls was
     adjusted to 0.1 mL/L. Each flask was inoculated with S. capricornutum to
     give an initial cell density of 5 x 102 cells/mL. The algal inoculum was
    prepared from an actively growing liquid culture of S. capricornutum in
     exponential growth phase.
* *
    Flasks were incubated in a temperature controlled orbital incubator under
     constant illumination (approximately 3000 lux) at 24±2°C for 4 days.
    Cell counts were made on days 2 and 4 using an electronic particle
     counter (Coulter counter). The temperature in the incubator was measured
    at 4-h intervals. The pH of the control and highest test concentration
    was measured on days 0, 2, and 4. Temperature remained within the 24±2°C
     specified range, and the pH ranged from 8.3 to 8.5 in the measured
     vessels.
* *
    All determination of EC50 values were based on nominal test
     concentrations and cell counts.
F020 243498
EOR
F002 455
F010 4.3
F004 3
F005 RE
F006 Stephenson R (1983). Isopropyl ether: Acute toxicity to Daphnia magna and
     Selenastrum capricornutum. Report # SBGR.83.215, Shell Research Ltd.
     Sittingbourne Research Centre, Sittingbourne, Kent, England.
F007 Stephenson R (1983). Isopropyl ether: Acute toxicity to Daphnia magna and
     Selenastrum capricornutum. Report # SBGR.83.215, Shell Research Ltd.
     Sittingbourne Research Centre, Sittingbourne, Kent, England.
F020 243502
EOR
F002 455
F010 4.3
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F004 3
F005 RL
F006 Test concentrations were not measured and there is no indication in the
     report whether the test vessels were sealed. The reported LC50 value may
     reflect a loss of test substance by volatilization if the flasks were not
     tightly sealed.
F007 Test concentrations were not measured and there is no indication in the
     report whether the test vessels were sealed. The reported LC50 value may
     reflect a loss of test substance by volatilization if the flasks were not
     tightly sealed.
F020 243501
EOR
F002 455
F010 4.3
F004 3
F005 RS
F006 96-hour EC50 = >1000 mg/L based on nominal concentrations.
* *
     The 96-hour cell counts in the treated flasks as a percent of the mean
     control cell counts were:
* *
      1.0 \text{ mg/L} = 84\%
                          46 \text{ mg/L} = 127\%
* *
      2.2 \text{ mg/L} = 108\%
                          100 \text{ mg/L} = 130\%
* *
      4.6 \text{ mg/L} = 91\%
                          22
F007 96-hour EC50 = >1000 mg/L based on nominal concentrations.
* *
     The 96-hour cell counts in the treated flasks as a percent of the mean
     control cell counts were:
* *
      1.0 \text{ mg/L} = 84\%
                          46 \text{ mg/L} = 127\%
* *
      2.2 \text{ mg/L} = 108\%
                          100 \text{ mg/L} = 130\%
* *
      4.6 \text{ mg/L} = 91\%
                          220 \text{ mg/L} = 113\%
* *
      10 \text{ mg/L} = 122\%
                          460 \text{ mg/L} = 127\%
* *
      22 \text{ mg/L} = 129\%
                          1000 \text{ mg/L} = 91\%
F020 243499
EOR
F002 455
F010 5.1.1
F004 5
F005 CL
F006 DIPE, when administered to adult male Sprague-Dawley rats, had an acute
     oral LD50 of >10 g/kg. 14-day immature rats were considerable more
     sensitive [LD50 4.5 g/kg].
F007 DIPE, when administered to adult male Sprague-Dawley rats, had an acute
     oral LD50 of >10 g/kg. 14-day immature rats were considerable more
     sensitive [LD50 4.5 g/kg].
F020 241249
EOR
F002 455
F010 5.1.1
F004 5
F005 ME
F006 Administered orally to nonfasted rats. LD50 calculated by the method of
     Litchfield and Wilcoxon [1949]. Similar to OECD 401.
F007 Administered orally to nonfasted rats. LD50 calculated by the method of
     Litchfield and Wilcoxon [1949]. Similar to OECD 401.
F020 241245
EOR
F002 455
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F010 5.1.1
F004 5
F005 RE
F006 Kimura ET, Ebert DM and Doge PW (1971). Acute toxicity and limits of
     solvent residues for sixteen organic solvents. Toxicol. Appl. Pharmacol.
     19:699-704.
F007 Kimura ET, Ebert DM and Doge PW (1971). Acute toxicity and limits of
     solvent residues for sixteen organic solvents. Toxicol. Appl. Pharmacol.
     19:699-704.
F020 241250
EOR
F002 455
F010 5.1.1
F004 5
F005 RL
F006 Not GLP but conducted at a reputable laboratory [Abbot Laboratories,
     Chicago].
F007 Not GLP but conducted at a reputable laboratory [Abbot Laboratories,
F020 241251
EOR
F002 455
F010 5.1.1
F004 5
F005 RM
F006 Test type: Acute oral toxicity
    Year: Prior to 1971
* *
    No. of animals/dose: 6 male for young adult and older adult
     6 - 12 male and female for 14-day old rats
* *
     Route of administration: Oral gavage
    Dose level: Variable
* *
* *
    Dose volume: Variable
F007 Test type: Acute oral toxicity
* *
     Year: Prior to 1971
    No. of animals/dose: 6 male for young adult and older adult
* *
    6 - 12 male and female for 14-day old rats
* *
    Route of administration: Oral gavage
* *
    Dose level: Variable
    Dose volume: Variable
* *
    Control group included: No, but none needed
F020 241248
EOR
F002 455
F010 5.1.1
F004 5
F005 RS
F006 14-day old: LD50 6.4 ml/kg [approx 4.5 g/kg]
     young adults: LD50 16.5 ml/kg [approx 11.6 g/kg]
* *
     Older adults: LD50 16.0 ml/kg [approx 11.2 g/kg]
* *
* *
     G/kg dose based on a density of 0.72 g/ml
F007 14-day old: LD50 6.4 ml/kg [approx 4.5 g/kg]
* *
     young adults: LD50 16.5 ml/kg [approx 11.6 g/kg]
* *
     Older adults: LD50 16.0 ml/kg [approx 11.2 g/kg]
* *
* *
     G/kg dose based on a density of 0.72 g/ml
F020 241247
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EOR
F002 455
F010 5.1.1
F004 5
F005 TC
F006 Rats were observed for up to 7 days after dosing.
F007 Rats were observed for up to 7 days after dosing.
F020 241246
EOR
F002 455
F010 5.1.1
F004 5
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Source/purity of test material is not specified, but stated to be
     analytical grade meeting ACS specifications.
F007 Diisopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Source/purity of test material is not specified, but stated to be
     analytical grade meeting ACS specifications.
F020 241244
EOR
F002 455
F010 5.1.1
F004 6
F005 CL
F006 The test article, when administered orally as received to New Zealand
     white rabbits had a minimal lethal dose of 7 - 9 ml/kg [approx 5 - 6.5
     g/kg].
F007 The test article, when administered orally as received to New Zealand
    white rabbits had a minimal lethal dose of 7 - 9 ml/kg [approx 5 - 6.5
     g/kg].
F020 241254
EOR
F002 455
F010 5.1.1
F004 6
F005 RE
F006 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
    Hyq. Toxicol. 21:72-96.
F007 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
     Hyg. Toxicol. 21:72-96.
F020 241256
EOR
F002 455
F010 5.1.1
F004 6
F005 RL
F006 Not conducted by GLP but at a reputable laboratory [Kettering Laboratory,
     University of Cincinnati].
F007 Not conducted by GLP but at a reputable laboratory [Kettering Laboratory,
    University of Cincinnati].
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F020 241255
EOR
F002 455
F010 5.1.1
F004 6
F005 RM
F006 Test type: Acute oral toxicity
    Year: Prior to 1939
* *
    Route of administration: Oral
* *
    Dose levels: 8.2, 7.2, 6.0, 5.2, 3.3, 1.62 g/kg
* *
    Dose volume: Variable
* *
    Control: No - none needed
F007 Test type: Acute oral toxicity
    Year: Prior to 1939
* *
    Route of administration: Oral
* *
    Dose levels: 8.2, 7.2, 6.0, 5.2, 3.3, 1.62 g/kg
* *
    Dose volume: Variable
* *
    Control: No - none needed
F020 241257
EOR
F002 455
F010 5.1.1
F004 6
F005 RS
F006 Minimal lethal dose between 7 - 9 ml/kg
* *
     The symptoms noted were lack of coordination and unsteadiness at onset
     followed by a slight narcosis. In the animals that died the narcosis
     progressed towards a deep narcosis with loss of corneal re
F007 Minimal lethal dose between 7 - 9 ml/kg
* *
* *
    The symptoms noted were lack of coordination and unsteadiness at onset
     followed by a slight narcosis. In the animals that died the narcosis
    progressed towards a deep narcosis with loss of corneal reflex and
    evidences of depressant action on the medulla appeared, respiration
    became progressively slower, irregular and variable in amplitude and drop
     in body temperature till respiration failed. In the surviving animals, no
     effect on HB, erythrocyte count, total and differential leukocyte count
    was observed. No delayed toxicity was observed during the recovery period
     of 4 months after treatment.
F020 241253
EOR
F002 455
F010 5.1.1
F004 6
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide nor added inhibitor.
F007 Diisopropyl ether (CAS No. 108-20-3)
    Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
    Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide nor added inhibitor.
F020 241252
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EOR
F002 455
F010 5.1.2
F004 2
F005 CL
F006 The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in
     three species.
F007 The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in
     three species.
F020 241267
EOR
F002 455
F010 5.1.2
F004 2
F005 RE
F006 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
     Hyg. Toxicol. 21:72-96.
F007 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
     Hyg. Toxicol. 21:72-96.
F020 241273
EOR
F002 455
F010 5.1.2
F004 2
F005 RL
F006 Not conducted by GLP. Few animals per group, but at a reputable
     laboratory [Kettering Laboratory, University of Cincinnati].
F007 Not conducted by GLP. Few animals per group, but at a reputable
     laboratory [Kettering Laboratory, University of Cincinnati].
F020 241272
EOR
F002 455
F010 5.1.2
F004 2
F005 RM
F006 Test type: Acute inhalation toxicity
     Year: Prior to 1939
* *
     No. animlas/sex/group: One to two animals per dose
* *
    Route of administration: Inhalation
* *
    Dose level: 0.3%; 1%; 3%; 6% in air
* *
    Dose volume: N/A
* *
     Control: No
F007 Test type: Acute inhalation toxicity
     Year: Prior to 1939
* *
     No. animlas/sex/group: One to two animals per dose
* *
    Route of administration: Inhalation
* *
    Dose level: 0.3%; 1%; 3%; 6% in air
* *
    Dose volume: N/A
* *
     Control: No
F020 241278
EOR
F002 455
F010 5.1.2
F004 2
F005 RS
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F006 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action
     1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia
     6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr
F007 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action
     1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia
     6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr
F020 241264
EOR
F002 455
F010 5.1.2
F004 2
F005 TC
F006 1 or 2 hrs or until death [6%]
F007 1 or 2 hrs or until death [6%]
F020 241259
EOR
F002 455
F010 5.1.2
F004 2
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Test material is stated to be commercial grade with 3% of isopropyl
* *
     alcohol, but with no peroxide or added inhibitor.
F007 Diisopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
    Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide or added inhibitor.
F020 241258
EOR
F002 455
F010 5.1.2
F004 3
F005 CL
F006 The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in
     three species.
F007 The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in
     three species.
F020 241268
EOR
F002 455
F010 5.1.2
F004 3
F005 RE
F006 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
     Hyg. Toxicol. 21:72-96.
F007 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
    Hyg. Toxicol. 21:72-96.
F020 241274
EOR
F002 455
F010 5.1.2
F004 3
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```
F005 RL
F006 Not conducted by GLP. Few animals per group, but at a reputable
     laboratory [Kettering Laboratory, University of Cincinnati].
F007 Not conducted by GLP. Few animals per group, but at a reputable
     laboratory [Kettering Laboratory, University of Cincinnati].
F020 241271
EOR
F002 455
F010 5.1.2
F004 3
F005 RM
F006 Test type: Acute inhalation toxicity
     Year: Prior to 1939
* *
     No. animlas/sex/group: One to two animals per dose
    Route of administration: Inhalation
* *
    Dose level: 0.3%; 1%; 3%; 6% in air
* *
    Dose volume: N/A
* *
     Control: No
F007 Test type: Acute inhalation toxicity
* *
    Year: Prior to 1939
* *
    No. animlas/sex/group: One to two animals per dose
* *
    Route of administration: Inhalation
* *
    Dose level: 0.3%; 1%; 3%; 6% in air
* *
    Dose volume: N/A
    Control: No
F020 241277
EOR
F002 455
F010 5.1.2
F004 3
F005 RS
F006 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action
     1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia
     6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr
F007 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action
     1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia
     6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr
F020 241265
EOR
F002 455
F010 5.1.2
F004 3
F005 TC
F006 1 or 2 hrs or until death [6%]
F007 1 or 2 hrs or until death [6%]
F020 241261
EOR
F002 455
F010 5.1.2
F004 3
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide or added inhibitor.
F007 Diisopropyl ether (CAS No. 108-20-3)
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Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
    Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide or added inhibitor.
F020 241260
EOR
F002 455
F010 5.1.2
F004 4
F005 CL
F006 The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in
     three species.
F007 The lowest lethal dose for 1 hr exposure was greater than 30,000 ppm in
     three species.
F020 241269
EOR
F002 455
F010 5.1.2
F004 4
F005 RE
F006 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
    Hyq. Toxicol. 21:72-96.
F007 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
     Hyg. Toxicol. 21:72-96.
F020 241275
EOR
F002 455
F010 5.1.2
F004 4
F005 RL
F006 Not conducted by GLP. Few animals per group, but at a reputable
     laboratory [Kettering Laboratory, University of Cincinnati].
F007 Not conducted by GLP. Few animals per group, but at a reputable
     laboratory [Kettering Laboratory, University of Cincinnati].
F020 241270
EOR
F002 455
F010 5.1.2
F004 4
F005 RM
F006 Test type: Acute inhalation toxicity
     Year: Prior to 1939
* *
    No. animlas/sex/group: One to two animals per dose
    Route of administration: Inhalation
* *
    Dose level: 0.3%; 1%; 3%; 6% in air
* *
    Dose volume: N/A
* *
    Control: No
F007 Test type: Acute inhalation toxicity
* *
    Year: Prior to 1939
* *
    No. animlas/sex/group: One to two animals per dose
* *
    Route of administration: Inhalation
* *
    Dose level: 0.3%; 1%; 3%; 6% in air
* *
    Dose volume: N/A
* *
    Control: No
F020 241276
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EOR
F002 455
F010 5.1.2
F004 4
F005 RS
F006 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action
     1 and 3.0 % (~30000 ppm)- 1 h: Not lethal; signs of anesthesia
     6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr
F007 0.3 % (~3000 ppm)- 2 h: No visible indication of anesthetic action
     1 and 3.0 % (\sim30000 ppm)- 1 h: Not lethal; signs of anesthesia
     6.0% (~60000 ppm): Death as the result of respiratory failure within 1 hr
F020 241266
EOR
F002 455
F010 5.1.2
F004 4
F005 TC
F006 1 or 2 hrs or until death [6%]
F007 1 or 2 hrs or until death [6%]
F020 241263
EOR
F002 455
F010 5.1.2
F004 4
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
    Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
    Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide or added inhibitor.
F007 Diisopropyl ether (CAS No. 108-20-3)
    Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
    Test material is stated to be commercial grade with 3% of isopropyl
    alcohol, but with no peroxide or added inhibitor.
F020 241262
EOR
F002 455
F010 5.1.3
F004 2
F005 CL
F006 The test article, when administered dermally to New Zealand white rabbits
     had an acute dermal LD50 of greater than 2.0 g/kg.
F007 The test article, when administered dermally to New Zealand white rabbits
     had an acute dermal LD50 of greater than 2.0 g/kg.
F020 241282
EOR
F002 455
F010 5.1.3
F004 2
F005 RE
F006 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
    Hyq. Toxicol. 21:72-96.
F007 Machle W, Scott EW and Treon J (1939). The physiological response to
     isopropyl ether and to a mixture of isopropyl ether and gasoline. J. Ind.
    Hyg. Toxicol. 21:72-96.
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F020 241284
EOR
F002 455
F010 5.1.3
F004 2
F005 RL
F006 Not GLP but conducted at a reputable laboratory [Kettering Laboratory,
     University of Cincinnati].
F007 Not GLP but conducted at a reputable laboratory [Kettering Laboratory,
     University of Cincinnati].
F020 241283
EOR
F002 455
F010 5.1.3
F004 2
F005 RM
F006 Test type: Acute dermal toxicity
     Year: Prior to 1939
* *
    No. of animals/sex/group: Unspecified
* *
    Route of administration: Dermal
* *
    Dose level: variable
* *
    Control: No
F007 Test type: Acute dermal toxicity
* *
    Year: Prior to 1939
    No. of animals/sex/group: Unspecified
* *
    Route of administration: Dermal
* *
    Dose level: variable
* *
     Control: No
F020 241285
EOR
F002 455
F010 5.1.3
F004 2
F005 RS
F006 No deaths or systemic effects were reported. In rabbits dermal unoccluded
     LD50 > 2.0 g/kg. The actual dose applied was much higher, but continued
     to evaporate from the skin during application.
F007 No deaths or systemic effects were reported. In rabbits dermal unoccluded
     LD50 > 2.0 g/kg. The actual dose applied was much higher, but continued
     to evaporate from the skin during application.
F020 241281
EOR
F002 455
F010 5.1.3
F004 2
F005 TC
F006 The material was continuously dripped onto the shaved skin to keep it wet
     for one hour, while continuously evaporating. 150 ml of material was used.
F007 The material was continuously dripped onto the shaved skin to keep it wet
     for one hour, while continuously evaporating. 150 ml of material was used.
F020 241280
EOR
F002 455
F010 5.1.3
F004 2
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
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```
Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide or added inhibitor.
F007 Diisopropyl ether (CAS No. 108-20-3)
* *
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Test material is stated to be commercial grade with 3% of isopropyl
     alcohol, but with no peroxide or added inhibitor.
F020 241279
EOR
F002 455
F010 5.11
F004 2
F005 CL
F006 DIPE does not appear to be a sensory irritant at concentrations up to 500
     ppm, but it does have an unpleasant odor at this concentration.
F007 DIPE does not appear to be a sensory irritant at concentrations up to 500
     ppm, but it does have an unpleasant odor at this concentration.
F020 241327
EOR
F002 455
F010 5.11
F004 2
F005 ME
F006 Non-quideline.
F007 Non-quideline.
F020 241324
EOR
F002 455
F010 5.11
F004 2
F005 RE
F006 Silverman LH, Schulte F and First MW (1946). Further studies on sensory
     response to certain industrial solvent vapors. J. Ind. Hyg. Toxicol.
     28(6):262-266.
F007 Silverman LH, Schulte F and First MW (1946). Further studies on sensory
     response to certain industrial solvent vapors. J. Ind. Hyg. Toxicol.
     28(6):262-266.
F020 241329
EOR
F002 455
F010 5.11
F004 2
F005 RL
F006 Not GLP but conducted at a reputable laboratory [Harvard School of Public
     Health, Boston].
F007 Not GLP but conducted at a reputable laboratory [Harvard School of Public
     Health, Boston].
F020 241328
EOR
F002 455
F010 5.11
F004 2
F005 RM
F006 Species/strain: Humans
** Sex: Male and female
```

```
Number/sex/group: Average of 12
* *
    Route of administration: Inhalation
* *
    Vehicle: None
* *
    Control: No
* *
    Year: Prior to 1946
* *
     GLP: No
F007 Species/strain: Humans
     Sex: Male and female
* *
    Number/sex/group: Average of 12
* *
    Route of administration: Inhalation
* *
    Vehicle: None
* *
    Control: No
* *
    Year: Prior to 1946
* *
    GLP: No
F020 241330
EOR
F002 455
F010 5.11
F004 2
F005 RS
F006 300 ppm: 35% of the subjects objected to this solvent because of the
     unpleasant odor rather than irritation.
     500 ppm: there was a sensory response that was acceptable to the
     majority of subjects.
F007 300 ppm: 35% of the subjects objected to this solvent because of the
     unpleasant odor rather than irritation.
* *
     500 ppm: there was a sensory response that was acceptable to the
     majority of subjects.
F020 241326
EOR
F002 455
F010 5.11
F004 2
F005 TC
F006 Subjects were exposed for 15 minutes and olfactory fatigue and irritation
     of mucous membranes were reported. "Motion pictures were shown to occupy
     the subject's attention and divert their thoughts from the atmospheric
     contamination to which
F007 Subjects were exposed for 15 minutes and olfactory fatigue and irritation
     of mucous membranes were reported. "Motion pictures were shown to occupy
     the subject's attention and divert their thoughts from the atmospheric
     contamination to which they were exposed."
F020 241325
EOR
F002 455
F010 5.11
F004 2
F006 Diisopropyl ether (CAS No. 108-20-3)
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Test material is stated to be technical grade product.
F007 Diisopropyl ether (CAS No. 108-20-3)
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Test material is stated to be technical grade product.
F020 241323
```

```
EOR
F002 455
F010 5.11
F004 3
F005 CL
F006 400 ppm: 5 mins of inhalation exposure caused no eye irritation, none to
     slight nose irritation, no pulmonary discomfort, olfactory recognition
     but no central nervous system effects.
* *
     800 ppm: 5 mins of inhalation exposed caused slight eye
F007 400 ppm: 5 mins of inhalation exposure caused no eye irritation, none to
     slight nose irritation, no pulmonary discomfort, olfactory recognition
     but no central nervous system effects.
* *
     800 ppm: 5 mins of inhalation exposed caused slight eye and nose
     irritation, none to slight pulmonary discomfort, definite olfactory
     recognition but no central nervous system effects.
F020 241335
EOR
F002 455
F010 5.11
F004 3
F005 ME
F006 Non-Guideline.
F007 Non-Guideline.
F020 241332
EOR
F002 455
F010 5.11
F004 3
F005 RE
F006 Hine CH, Anderson HH and Kodama JK (1955). Sensory thresholds of certain
     Shell organic solvents, Progress Report 1, Report to Shell Development
     Company, November 15, 1955. UC Report No. 247.
F007 Hine CH, Anderson HH and Kodama JK (1955). Sensory thresholds of certain
     Shell organic solvents, Progress Report 1, Report to Shell Development
     Company, November 15, 1955. UC Report No. 247.
F020 241337
EOR
F002 455
F010 5.11
F004 3
F005 RL
F006 Not GLP but conducted at a reputable laboratory [University of California
     School of Medicine].
F007 Not GLP but conducted at a reputable laboratory [University of California
     School of Medicine].
F020 241336
EOR
F002 455
F010 5.11
F004 3
F005 RM
F006 Species/strain: Young adult humans [University of California staff and
    medical students]
* *
     Sex: Not specified
     Number/sex/group: Not specified
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Route of administration: Inhalation
    Vehicle: None
* *
    Control: No
* *
    Year: 1955
* *
     GLP: No
F007 Species/strain: Young adult humans [University of California staff and
    medical students
     Sex: Not specified
* *
    Number/sex/group: Not specified
* *
    Route of administration: Inhalation
* *
    Vehicle: None
* *
    Control: No
* *
    Year: 1955
* *
    GLP: No
F020 241338
EOR
F002 455
F010 5.11
F004 3
F005 RS
F006 Numbers of subjects with degree of effect
                           400 ppm
     Concentration
                                              mqq 008
* *
    Number subjects:
                                              7
* *
                           7 absent
    Eye irritation:
                                        3 absent, 3 slight, 1 mod.
    Nose irritation: 5 absent, 2 slight
                                              2 abse
F007 Numbers of subjects with degree of effect
* *
    Concentration
                           400 ppm
                                              mqq 008
* *
    Number subjects:
                           7 absent
    Eye irritation:
                                        3 absent, 3 slight, 1 mod.
    Nose irritation: 5 absent, 2 slight
* *
                                           2 absent, 5 slight
* *
    Pulmonary discomfort: 7 absent
                                              4 absent, 3 slight
* *
    Olfactory cognition: 1 slight, 6 mod. 4 mod., 3 severe
* *
    CNS effects :
                                             7 absent
                          7 absent
F020 241334
EOR
F002 455
F010 5.11
F004 3
F005 TC
F006 Exposures were conducted in a whole-body chamber approximately 7700 l
     equipped with a fan. Exposures were made in a static atmosphere generated
    by vaporizing a predetermined quantity of test solvent from a hot
     surface. Five minutes were all
F007 Exposures were conducted in a whole-body chamber approximately 7700 l
     equipped with a fan. Exposures were made in a static atmosphere generated
    by vaporizing a predetermined quantity of test solvent from a hot
     surface. Five minutes were allowed for evaporation and equilibration, and
     subjects were exposed for 5 minutes, during which time they noted the
     degree of subjective responses at one-minute intervals.
F020 241333
EOR
F002 455
F010 5.11
F004 3
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
    Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
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[propane], 2,2 '-
     Test material is stated to be commercial grade with purity of 98% or
     better, provided by Shell Chemical Corporation.
F007 Diisopropyl ether (CAS No. 108-20-3)
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Test material is stated to be commercial grade with purity of 98% or
     better, provided by Shell Chemical Corporation.
F020 241331
EOR
F002 455
F010 5.3
F004 2
F005 CL
F006 Di-isopropanol was negative in this in vitro assay.
F007 Di-isopropanol was negative in this in vitro assay.
F020 241289
EOR
F002 455
F010 5.3
F004 2
F005 RE
F006 Wass U and Belin L (1990). An in vitro method for predicting sensitizing
     properties of inhaled chemicals. Scand. J. Work Environ. Health.
     116:208-214.
F007 Wass U and Belin L (1990). An in vitro method for predicting sensitizing
     properties of inhaled chemicals. Scand. J. Work Environ. Health.
     116:208-214.
F020 241291
EOR
F002 455
F010 5.3
F004 2
F005 RL
F006 Not conducted by GLP; research method not accepted by regulatory
     agencies; in vitro surrogate for respiratory sensitisation.
F007 Not conducted by GLP; research method not accepted by regulatory
     agencies; in vitro surrogate for respiratory sensitisation.
F020 241290
EOR
F002 455
F010 5.3
F004 2
F005 RM
F006 Route of administration: N/A
    Sex: N/A
* *
     Dose level: N/A
* *
    Dose volume: N/A
* *
     Control group included: Positive and negative controls included
F007 Route of administration: N/A
* *
     Sex: N/A
* *
    Dose level: N/A
    Dose volume: N/A
* *
     Control group included: Positive and negative controls included
F020 241292
EOR
F002 455
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```
F010 5.3
F004 2
F005 RS
F006 Diisopropanol was negative in this in vitro assay for potential
     respiratory sensitization. The assay gave positive responses with
     several known respiratory sensitizers.
F007 Diisopropanol was negative in this in vitro assay for potential
    respiratory sensitization. The assay gave positive responses with
     several known respiratory sensitizers.
F020 241288
EOR
F002 455
F010 5.3
F004 2
F005 TC
F006 A method for monitoring chemical reactivity in aqueous solutions, at
     neutral pH and 37 degrees C, was developed. The chemical was allowed to
    react with a lysine-containing peptide, and the reaction was monitored
     with high-performance liquid
F007 A method for monitoring chemical reactivity in aqueous solutions, at
    neutral pH and 37 degrees C, was developed. The chemical was allowed to
    react with a lysine-containing peptide, and the reaction was monitored
    with high-performance liquid chromatography. Simple acids, bases, and
     solvents did not react with the peptide, whereas isocyanates, anhydrides,
    and chloramine-T, substances well known for their sensitizing and asthma
     inducing properties, did. Thus a positive test strongly suggested that
     the chemical had the potential to act as a hapten and cause sensitization
     when inhaled.
F020 241287
EOR
F002 455
F010 5.3
F004 2
F005 TS
F006 Diisopropyl ether (CAS No.108-20-3)
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Source/purity not specified.
F007 Diisopropyl ether (CAS No.108-20-3)
    Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Source/purity not specified.
F020 241286
EOR
F002 455
F010 5.4
F004 5
F005 CL
F006 NOAEL = 480 \text{ ppm}
F007 NOAEL = 480 ppm
F020 241297
EOR
F002 455
F010 5.4
F004 5
F005 ME
F006 Statistical method:
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Statistical analyses of numerical data included ANOVA and Tukey's
     studentized range test for data on serum chemistry. Duncan's multiple
     range test was used for hematology and body weights to assess
     statistically signific
F007 Statistical method:
     Statistical analyses of numerical data included ANOVA and Tukey's
     studentized range test for data on serum chemistry. Duncan's multiple
    range test was used for hematology and body weights to assess
     statistically significant differences between control and exposed groups.
F020 241294
EOR
F002 455
F010 5.4
F004 5
F005 RE
F006 Dalbey W and Fueston M (1996). Subchronic and Developmental Toxicity
     Studies of Vaporized Diisopropyl Ether in Rats. Journal of Toxicology and
     Environmental Health 49:29-43.
F007 Dalbey W and Fueston M (1996). Subchronic and Developmental Toxicity
     Studies of Vaporized Diisopropyl Ether in Rats. Journal of Toxicology and
     Environmental Health 49:29-43.
F020 241299
EOR
F002 455
F010 5.4
F004 5
F005 RL
F006 GLP unknown. Study well documented, meets generally accepted scientific
     principles, acceptable for assessment.
F007 GLP unknown. Study well documented, meets generally accepted scientific
     principles, acceptable for assessment.
F020 241298
EOR
F002 455
F010 5.4
F004 5
F005 RM
F006 Male and female rats were acclimated for 2 weeks before initiation of
     exposures that began at ~8 weeks of age. Exposed animals were
     individually housed in 1-m3 inhalation chambers. Untreated control
     animals were housed in a separate room in
F007 Male and female rats were acclimated for 2 weeks before initiation of
     exposures that began at ~8 weeks of age. Exposed animals were
     individually housed in 1-m3 inhalation chambers. Untreated control
     animals were housed in a separate room in identical caging. Room
     environment was set to 20-22°C and 40-60% relative humidity. Lights were
     on a 12/12-hr light/dark cycle. Food and water were provided ad libitum
     except during exposures.
* *
    Vapors were generated by metering DIPE on to a warmed fiberglass wick and
     carried to the three 1m3 exposure chambers by a stream of measured and
     filtered room air. Temperature and humidity in the chambers were measured
```

every 30 minutes and air flow through each chamber was metered to provide at least 12 air changes/hour. Chamber air samples were collected with a gas-tight syringe and analyzed with a gas chromatograph with a flame ionization detector and a fused silica column. Samples were periodically

obtained for analysis by gas chromatography/ mass spectroscopy.

* * The primary endpoints during the course of exposures were individual body weights recorded weekly and clinical signs recorded daily except on weekends. * * Following the last exposure, rats were fasted overnight and weighed; blood samples were obtained via the orbital sinus using light anesthesia. Samples were used to determine values for WBC, RBC, Hgb, Hct, MCV, MCH, MCHC, platelets, and differential counts. Glucose, urea nitrogen, total protein, albumin, globulin, A/G ratio, sorbitol dehydrogenase, aspartate aminotransferase, alanine aminotransferase, alkaline phosphatase, total bilirubin, creatinine, cholesterol, triglycerides, uric acid, Cl, Ca, Na, K, and P. Following the collection of blood samples, all animals were euthanized with sodium pentobarbital (i.p.) and exanguinated. Approximately 40 tissues were collected for histopathology and organs were weighed. Testis and associated tissues were preserved whole in 10% buffered formalin except for the left cauda epididymis of 10 rats in both control groups and the highest test group; epididymides were evaluated for morphology and number of sperm. The left testis in these groups was weighed and used for determination of number of testicular spermatids. F020 241296 EOR F002 455 F010 5.4 F004 5 F005 RM F006 Type: 90-Day Subchronic Species/strain: Rat/Sprague-Dawley; Tac:N(SD)fBR No./sex/dose: 14/sex/group * * Vehicle: None * * Method: USEPA 1984, 40CFR Part 798:2450 F007 Type: 90-Day Subchronic * * Species/strain: Rat/Sprague-Dawley; Tac:N(SD)fBR * * No./sex/dose: 14/sex/group * * Vehicle: None Method: USEPA 1984, 40CFR Part 798:2450 F020 241300 EOR F002 455 F010 5.4 F004 5 F005 RS F006 DIPE did not adversely affect clinical signs body weight, serum chemistry, hematology, or the number of sperm or spermatids. Exposure to males at 7100 ppm resulted in hypertrophy of liver cells associated with increased liver weight and in F007 DIPE did not adversely affect clinical signs body weight, serum chemistry, hematology, or the number of sperm or spermatids. Exposure to males at 7100 ppm resulted in hypertrophy of liver cells associated with increased liver weight and in increased kidney weight with an increased incidence of hyaline droplets in proximal tubules of the kidney. Females had increased weight of both liver and kidney, although kidney increased only in relation to sham-exposed controls and no morphologic changes were observed in either organ. At 3300 ppm, weights of liver and kidney were increased in males; the liver weights were increased in females only compared to sham-exposed controls and not untreated controls. No

morphologic abnormalities were observed. No changes were observed with

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480 ppm.
F020 241295
EOR
F002 455
F010 5.4
F004 5
F005 TS
F006 Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.
F007 Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.
F020 241293
EOR
F002 455
F010 5.4
F004 6
F005 CL
F006 Inhalation exposures to DIPE at concentrations as high as 7060 ppm for 13
     weeks resulted in few observable effects on the nervous system.
F007 Inhalation exposures to DIPE at concentrations as high as 7060 ppm for 13
     weeks resulted in few observable effects on the nervous system.
F020 241305
EOR
F002 455
F010 5.4
F004 6
F005 ME
F006 Statistical method:
    All statistical analyses were performed with SAS software.
    rectal temperatures fore- and hindlimb grip strengths, the number of
     rears, and motor activity were analyzed by a one-way analysis of variance
     fol
F007 Statistical method:
    All statistical analyses were performed with SAS software. Body weights,
    rectal temperatures fore- and hindlimb grip strengths, the number of
    rears, and motor activity were analyzed by a one-way analysis of variance
     followed by Duncan's multiple range test. The remaining data from the FOB
    were analyzed by Fisher's exact test using an extended contingency table
    containing all four groups of at given sex at a specified time. If a
     significant difference occurred for a given parameter, Fisher's exact
    test was used to directly compare each group individually against the
     control. Brain weights, lengths and widths, were analyzed by Student's
     t-test.
F020 241302
E \cap R
F002 455
F010 5.4
F004 6
F005 RE
F006 Rodriguez SC and Dalbey W (1997). Subchronic neurotoxicity of vaporized
     Diisopropyl Ether in rats. International Journal of Toxicology 16:599-610.
F007 Rodriguez SC and Dalbey W (1997). Subchronic neurotoxicity of vaporized
     Diisopropyl Ether in rats. International Journal of Toxicology 16:599-610.
F020 241307
EOR
F002 455
F010 5.4
F004 6
F005 RL
```

F006 GLP unknown. Study well documented, meets generally accepted scientific principles, acceptable for assessment. F007 GLP unknown. Study well documented, meets generally accepted scientific principles, acceptable for assessment. F020 241306 EOR F002 455 F010 5.4 F004 6 F005 RM F006 Male and female rats were acclimated for 2 weeks before initiation of exposures that began at ~8 weeks of age. Sham-exposed and exposed animals were individually housed in 1-m3 inhalation chambers except during behavioral testing, when they F007 Male and female rats were acclimated for 2 weeks before initiation of exposures that began at ~8 weeks of age. Sham-exposed and exposed animals were individually housed in 1-m3 inhalation chambers except during behavioral testing, when they were placed in another room overnight and evaluated the following day. Room environment was set to 20-22°C and 40-60% relative humidity. Lights were on a 12/12-hr light/dark cycle. Food and water were provided ad libitum except during exposures. Exposure vapors were generated by metering DIPE on to a warmed fiberglass wick and carried to the three 1-m3 exposure chambers by a stream of measured and filtered room air. Temperature and humidity in the chambers were measured every 30 minutes and air flow through each chamber was metered to provide at least 12 air changes/hour. Chamber air samples were collected with a gas-tight syringe and analyzed with a gas chromatograph with a flame ionization detector and a fused silica column. Samples were periodically obtained for analysis by gas chromatography/ mass spectroscopy. * * Exposures were stagger-started over a 5-day period with 16 animals, 2/sex/group, receiving their first exposure on each of 5 consecutive * * The rats were observed for signs of toxicity daily prior to initiation of exposures, and individual body weights were recorded weekly. * * During weeks of Functional Observation Battery (FOB) evaluation, four rats/group would be removed from the inhalation chamber and housed separately overnight and evaluated on the following day between the hours of 7:30 a.m. and 11:00 a.m. The process was repeated for each consecutive day until all rats were evaluated. The rats were evaluated with minimal disruption to the exposure schedule and with a manageable number of rats per day, 16 on any given day. The animals were evaluated in an FOB followed by a determination of motor activity prior to initiation of exposure; the FOB following 2, 4, 8, and 13 weeks of exposures, and for motor activity following 4, 8, and 13 weeks of exposures. Following the final determination of motor activity, the animals were anesthetized, intravascularly perfused, and the brain, spinal cord, and peripheral nerves removed for microscopic examination. The FOB consisted of initially observing home-cage positioning, posture, and reaction to removal from the cage. This was followed by evaluation for exophthalmus/palpebral closure, lacrimation, salivation, pupillary response, palpebral reflex, and pinna reflex. These observations were scored by type and intensity. The animals were then observed for open

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field behavior. Piloerection, respiratory rate, tremors, convulsions,
    posture, gait, ataxic gait, tail elevation, unperturbed activity level,
     vocalization, number of rears, fecal balls, and urine pools were all
     recorded during the open-field observations. Reactions to the approach of
     a pencil, finger snap, and tail pinch were ranked and recorded. Finally,
     fore- and hindlimb grip strength , rectal temperature , and body weight
     were measured. Automated motor activity was assessed for 30 minutes in
     figure-eight mazes after the completion of the FOB.
    Following the last FOB and motor activity evaluation, the rats were
    anesthetized with heparinized sodium pentobarbital (i.p.). The thoracic
     cavity was opened and the animals were infused with phosphate-buffered
    gluteraldehyde through the left ventricle. The perfused brain, spinal
     cord, and sciatic nerve with its tibial, sural, and peroneal divisions
    were removed. The brain and nerve tissues were processed for embedding in
    paraffin or glycol methacrylate (dorsal root ganglia and peripheral
     nerves) and sectioned for light or electron microscopic pathologic
     evaluation.
F020 241304
EOR
F002 455
F010 5.4
F004 6
F005 RM
F006 Type: 90-Day Neurotoxicity
     Species/strain: Rat/Sprague-Dawley; Tac:N(SD)fBR
* *
    No./sex/dose: 10/sex/group
* *
    Vehicle: None
* *
    Method: USEPA 1984, 40CFR Part 798:6050, 6400, and 6200
F007 Type: 90-Day Neurotoxicity
* *
     Species/strain: Rat/Sprague-Dawley; Tac:N(SD)fBR
* *
    No./sex/dose: 10/sex/group
* *
    Vehicle: None
* *
    Method: USEPA 1984, 40CFR Part 798:6050, 6400, and 6200
F020 241308
EOR
F002 455
F010 5.4
F004 6
F005 RS
F006 Motor activity in a figure-eight maze and unperturbed activity in the FOB
     were decreased at week 4 in females exposed to 7060 ppm; activity in the
     FOB was also decreased in females exposed to 450 ppm at week 4. Other
     changes in the FOB app
F007 Motor activity in a figure-eight maze and unperturbed activity in the F0B
     were decreased at week 4 in females exposed to 7060 ppm; activity in the
     FOB was also decreased in females exposed to 450 ppm at week 4. Other
     changes in the FOB appeared to be minor, and no changes were observed
     during microscopic examination of tissues from the nervous system.
F020 241303
EOR
F002 455
F010 5.4
F004 6
F005 TS
F006 Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.
F007 Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.
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F020 241301
EOR
F002 455
F010 5.5
F004 4
F005 CL
F006 Under the conditions of this study, the test material was not mutagenic.
F007 Under the conditions of this study, the test material was not mutagenic.
F020 241311
EOR
F002 455
F010 5.5
F004 4
F005 RE
F006 Brooks TM, Meyer AL and Hutson DH (1988). The genetic toxicology of some
     hydrocarbon and oxygenated solvents. Mutagenesis 3(3):227-232.
F007 Brooks TM, Meyer AL and Hutson DH (1988). The genetic toxicology of some
     hydrocarbon and oxygenated solvents. Mutagenesis 3(3):227-232.
F020 241313
EOR
F002 455
F010 5.5
F004 4
F005 RL
F006 Restriction due to the lack of any information regarding the selection of
     dose levels used during the study. In addition no information is
    presented regarding cytotoxicity or the presence of test material
    precipitate in the cultures. Altho
F007 Restriction due to the lack of any information regarding the selection of
     dose levels used during the study. In addition no information is
    presented regarding cytotoxicity or the presence of test material
    precipitate in the cultures. Although study was not stated to be
     conducted under GLP, it was conducted at a reputable laboratory [Shell
     Research Limited, Sittingbourne Research Center].
F020 241312
EOR
F002 455
F010 5.5
F004 4
F005 RM
F006 Strains tested: Salmonella typhimurium tester strains TA98, TA100,
     TA1535, TA1537, TA1538
* *
* *
     Exposure method: Preincubation assay for volatile compounds [Brooks and
    Dean 19811
* *
* *
     Test Substance Doses/concentration levels: Up to 8000 ug/ml i
F007 Strains tested: Salmonella typhimurium tester strains TA98, TA100,
     TA1535, TA1537, TA1538
* *
    Exposure method: Preincubation assay for volatile compounds [Brooks and
    Dean 1981]
* *
    Test Substance Doses/concentration levels: Up to 8000 ug/ml in the
    pre-incubation mix
* *
* *
    Metabolic activation: With and without (S9 fraction mix of livers of
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Aroclor 1254 pretreated rats)
* *
     Vehicle: Tween 80/ethanol
* *
* *
     Tester strain, activation status, Positive Controls and concentration
     level: Not stated
* *
* *
     Statistical analysis: Mean revertant colony count and standard deviation
     were determined for each dose point.
* *
* *
    Dose Rangefinding Study: Cytotoxicity study
* *
**
     S9 Optimization Study: No
F020 241314
E \cap R
F002 455
F010 5.5
F004 4
F005 RS
F006 DIPE did not induce reverse gene mutation in any strain. The test
     substance was not genotoxic in this assay with or without metabolic
     activation.
F007 DIPE did not induce reverse gene mutation in any strain. The test
     substance was not genotoxic in this assay with or without metabolic
     activation.
F020 241310
EOR
F002 455
F010 5.5
F004 4
F005 TS
F006 Diisopropyl ether (CAS No. 108-20-3)
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Source/purity not specified.
F007 Diisopropyl ether (CAS No. 108-20-3)
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Source/purity not specified.
F020 241309
EOR
F002 455
F010 5.5
F004 5
F005 CL
F006 Under the conditions of this study, the test material was not mutagenic.
F007 Under the conditions of this study, the test material was not mutagenic.
F020 242883
EOR
F002 455
F010 5.5
F004 5
F005 RE
F006 Brooks T.M., Meyer A.L. and Hutson D.H. (1988). The genetic toxicology
     of some hydrocarbon and oxygenated solvents. Mutagenesis, 3(3):227-232.
F007 Brooks T.M., Meyer A.L. and Hutson D.H. (1988). The genetic toxicology
     of some hydrocarbon and oxygenated solvents. Mutagenesis, 3(3):227-232.
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F020 242885
EOR
F002 455
F010 5.5
F004 5
F005 RL
F006 Restriction due to the lack of any information regarding the selection of
     dose levels used during the study. In addition no information is
     presented regarding cytotoxicity or the presence of test material
     precipitate in the cultures. Altho
F007 Restriction due to the lack of any information regarding the selection of
     dose levels used during the study. In addition no information is
    presented regarding cytotoxicity or the presence of test material
    precipitate in the cultures. Although study was not stated to be
     conducted under GLP, it was conducted at a reputable laboratory [Shell
     Research Limited, Sittingbourne Research Center].
F020 242884
EOR
F002 455
F010 5.5
F004 5
F005 RM
F006 Test type: Chromosome damage
* *
* *
     Exposure method: For volatile compounds
* *
* *
    Metabolic activation: Metabolic activation S9 was not added because
     liver cells are metabolically competent
* *
* *
    Vehicle: Tween 80/ethanol
* *
* *
     Tester strain, activation sta
F007 Test type: Chromosome damage
* *
* *
     Exposure method: For volatile compounds
* *
* *
    Metabolic activation: Metabolic activation S9 was not added because
    liver cells are metabolically competent
    Vehicle: Tween 80/ethanol
* *
* *
    Tester strain, activation status, Positive Controls and concentration
     level: Cultured CHO cells were grown in 80 cm2 flasks for 24 hr before
     compound treatment. Treatment periods were 5 hr in the presence of S9 mix
     and 24 hr in the absence of S9. Colcemid was added to all cultures 22 hr
     after the initial treatment. After a further 2 hr, the cells were
     trypsinized, resuspended in hypotonic solution and then fixed, before
     spotting onto slides. Cell preparations were then stained with Giemsa.
     The slides were randomly coded and 100 cells from each culture were
     analyzed microscopically. Mitotic index estimations were also made. The
    positive controls were ethylmethanesulfonate [-S9] and cyclophosphamide
     [+S9].
* *
    Vehicle control: Yes
* *
* *
    Dose rangefinding study: Cytotoxicity study
* *
```

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S9 Optimization study: No
F020 242886
EOR
F002 455
F010 5.5
F004 5
F005 RS
F006 DIPE did not induce chromosomal damage in CHO cells. The test substance
     was not genotoxic in this assay.
F007 DIPE did not induce chromosomal damage in CHO cells. The test substance
    was not genotoxic in this assay.
F020 242882
EOR
F002 455
F010 5.5
F004 5
F005 TS
F006 Di-isopropyl ether (CAS No. 108-20-3)
     Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Source/purity of test material: 98.5%
F007 Di-isopropyl ether (CAS No. 108-20-3)
    Chemical Name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Source/purity of test material: 98.5%
F020 242881
EOR
F002 455
F010 5.5
F004 6
F005 CL
F006 Under the conditions of this study, the test material was not mutagenic.
F007 Under the conditions of this study, the test material was not mutagenic.
F020 242889
EOR
F002 455
F010 5.5
F004 6
F005 RE
F006 Brooks T.M., Meyer A.L. and Hutson D.H. (1988). The genetic toxicology
     of some hydrocarbon and oxygenated solvents. Mutagenesis, 3(3):227-232.
F007 Brooks T.M., Meyer A.L. and Hutson D.H. (1988). The genetic toxicology
     of some hydrocarbon and oxygenated solvents. Mutagenesis, 3(3):227-232.
F020 242891
EOR
F002 455
F010 5.5
F004 6
F005 RL
F006 Restriction due to the lack of any information regarding the selection of
     dose levels used during the study. In addition no information is
    presented regarding cytotoxicity or the presence of test material
    precipitate in the cultures. Altho
F007 Restriction due to the lack of any information regarding the selection of
    dose levels used during the study. In addition no information is
    presented regarding cytotoxicity or the presence of test material
    precipitate in the cultures. Although study was not stated to be
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conducted under GLP, it was conducted at a reputable laboratory [Shell
     Research Limited, Sittingbourne Research Center].
F020 242890
EOR
F002 455
F010 5.5
F004 6
F005 RM
F006 Test type: Chromosome damage
* *
     Strains tested: RL4
* *
* *
     Metabolic activation: Metabolic activation S9 was not added because
     liver cells are metabolically competent.
* *
* *
     Vehicle: Tween 80/ethanol
* *
* *
     Tester strain, activation status, Positive Contr
F007 Test type: Chromosome damage
* *
* *
     Strains tested: RL4
* *
* *
     Metabolic activation: Metabolic activation S9 was not added because
     liver cells are metabolically competent.
* *
* *
     Vehicle: Tween 80/ethanol
* *
     Tester strain, activation status, Positive Controls and concentration
     level: Cultured rat liver cells were grown and treated on glass
     microscope slides contained in 100 ml glass Leighton tubes. After 22 hr
     exposure to test compound or solvent, colcemid was added to each culture.
     After a further 2 hr, the slides were removed, subjected to hypotonic
     treatment followed by fixation and stained with Giemsa. The preparations
     were randomly coded and 100 cells from each culture were analyzed
     microscopically. The positive control was 7,12-dimethylbenzanthracenene.
* *
     Vehicle control: Yes
* *
* *
    Dose rangefinding study: Cytotoxicity study
* *
* *
     S9 Optimization study: None needed
F020 242892
E \cap R
F002 455
F010 5.5
F004 6
F005 RS
F006 DIPE did not induce chromosomal damage in rat liver cells. The test
     substance was not genotoxic in this assay.
F007 DIPE did not induce chromosomal damage in rat liver cells. The test
     substance was not genotoxic in this assay.
F020 242888
EOR
F002 455
F010 5.5
F004 6
F005 TS
```

```
F006 Di-isopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
* *
     Source/purity of test material: 98.5%
F007 Di-isopropyl ether (CAS No. 108-20-3)
     Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Source/purity of test material: 98.5%
F020 242887
EOR
F002 455
F010 5.5
F004 7
F005 CL
F006 Under the conditions of this study, the test material was not genotoxic.
F007 Under the conditions of this study, the test material was not genotoxic.
F020 242895
EOR
F002 455
F010 5.5
F004 7
F005 RE
F006 Brooks T.M., Meyer A.L. and Hutson D.H. (1988). The genetic toxicology
     of some hydrocarbon and oxygenated solvents. Mutagenesis, 3(3):227-232.
F007 Brooks T.M., Meyer A.L. and Hutson D.H. (1988). The genetic toxicology
     of some hydrocarbon and oxygenated solvents. Mutagenesis, 3(3):227-232.
F020 242897
EOR
F002 455
F010 5.5
F004 7
F005 RL
F006 Restriction due to the lack of any information regarding the selection of
     dose levels used during the study. In addition no information is
     presented regarding cytotoxicity or the presence of test material
     precipitate in the cultures. Altho
F007 Restriction due to the lack of any information regarding the selection of
     dose levels used during the study. In addition no information is
     presented regarding cytotoxicity or the presence of test material
     precipitate in the cultures. Although study was not stated to be
     conducted under GLP, it was conducted at a reputable laboratory [Shell
     Research Limited, Sittingbourne Research Center].
F020 242896
EOR
F002 455
F010 5.5
F004 7
F006 Test type: Yeast mitotic gene conversion
* *
* *
     Strains tested: JD1
* *
* *
     Exposure method: [Brooks and Dean 1981]
* *
* *
     Metabolic activation: With and without (S9 fraction mix of livers of
     Aroclor 1254 pretreated rats)
```

```
* *
    Vehicle: Tween 80/ethanol
* *
* *
F007 Test type: Yeast mitotic gene conversion
* *
* *
     Strains tested: JD1
* *
* *
     Exposure method: [Brooks and Dean 1981]
* *
* *
    Metabolic activation: With and without (S9 fraction mix of livers of
    Aroclor 1254 pretreated rats)
* *
    Vehicle: Tween 80/ethanol
* *
    Tester strain, activation status, Positive Controls and concentration
     level: Yeast cells were grown in log-phase, washed and resuspended in
     2/5 strength YEPD broth at a concentration of 1 X 107 cells/ml. The
     suspension was divided into 1.9 ml amounts in 30 ml universal containers
     and 0.1 ml of test compound solution was added. For experiments with
    metabolic activation [+S9], 0.1 ml of DIPE was added to 01.6 ml of yeast
    cell suspension, together with 0.3 ml of S9 mix. Initially a range of
     concentrations of DIPE was tested up to 5 mg/ml if solubility allowed. A
     second experiment was performed based on these results and taking into
    account cell viability. The cultures were incubated with shaking at 30 C
     for 18 hr. Aliquots were plated onto the appropriate culture media for
     selection of mitotic gene convertants and cells surviving the treatment.
    Mitotic gene conversion may be scored by supplementing the minimal medium
    with histidine to score tryptophan prototrophs, and with tryptophan to
     score histidine prototrophs. Control plates were set up with solvent
     alone and with the positive control compounds 4-nitroquinoline oxide and
     cyclophosphamide.
* *
    Vehicle control: Yes
* *
* *
    Dose rangefinding study: Cytotoxicity study
* *
* *
     S9 Optimization study: No
F020 242898
EOR
F002 455
F010 5.5
F004 7
F005 RS
F006 DIPE did not induce mitotic gene conversion I yeast. The test substance
     was not genotoxic in this assay with or without metabolic activation.
F007 DIPE did not induce mitotic gene conversion I yeast. The test substance
    was not genotoxic in this assay with or without metabolic activation.
F020 242894
EOR
F002 455
F010 5.5
F004 7
F005 TS
F006 Di-isopropyl ether (CAS No. 108-20-3)
    Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Source/purity of test material: 98.5%
```

```
F007 Di-isopropyl ether (CAS No. 108-20-3)
    Chemical name: Isopropyl ether; isopropoxy propane, 2-; oxybis
     [propane], 2,2 '-
     Source/purity of test material: 98.5%
F020 242893
EOR
F002 455
F010 5.8.2
F004 2
F005 CL
F006 DIPE is not a teratogen.
F007 DIPE is not a teratogen.
F020 241319
EOR
F002 455
F010 5.8.2
F004 2
F005 ME
F006 Statistical method:
     Statistical analyses of numerical data included ANOVA and Tukey's
     studentized range test for data on serum chemistry. Duncan's multiple
    range test was used for hematology and body weights to assess
     statistically signific
F007 Statistical method:
    Statistical analyses of numerical data included ANOVA and Tukey's
     studentized range test for data on serum chemistry. Duncan's multiple
    range test was used for hematology and body weights to assess
     statistically significant differences between control and exposed groups.
    Data on the maternal biophase, cesarean sections, and fetuses were
     evaluated by ANOVA followed by group comparisons using Fisher's exact or
     Dunnett's test.
F020 241316
EOR
F002 455
F010 5.8.2
F004 2
F005 RE
F006 Dalbey W and Fueston M (1996). Subchronic and Developmental Toxicity
     Studies of Vaporized Diisopropyl Ether in Rats. Journal of Toxicology and
     Environmental Health 49:29-43.
F007 Dalbey W and Fueston M (1996). Subchronic and Developmental Toxicity
     Studies of Vaporized Diisopropyl Ether in Rats. Journal of Toxicology and
     Environmental Health 49:29-43.
F020 241321
EOR
F002 455
F010 5.8.2
F004 2
F005 RL
F006 GLP unknown. Study well documented, meets generally accepted scientific
     principles, acceptable for assessment.
F007 GLP unknown. Study well documented, meets generally accepted scientific
    principles, acceptable for assessment.
F020 241320
EOR
F002 455
F010 5.8.2
```

F004 2
F005 RM
F006 Nulliparous females were housed with males in a 1:1 ratio and observed
* daily for evidence of breeding activity. Females positive for sperm plug
* and for sperm in the vaginal lavage fluid were considered to be at day 0
* of gestation and were i
F007 Nulliparous females were housed with males in a 1:1 ratio and observed
* daily for evidence of breeding activity. Females positive for sperm plug
* and for sperm in the vaginal lavage fluid were considered to be at day 0
* of gestation and were individually housed. The females were then randomly

distributed to 5 groups of 22 animals each: untreated controls, sham-exposed controls, and 3 groups exposed to vapors of DIDP at 430, 3095, or 6745 ppm for 6 hr/day on gestation days (GD) 6-15.

** Vapors were generated by metering DIPE on to a warmed fiberglass wick and carried to the three 1m3 exposure chambers by a stream of measured and filtered room air. Temperature and humidity in the chambers were measured every 30 minutes and air flow through each chamber was metered to provide at least 12 air changes/hour. Chamber air samples were collected with a gas-tight syringe and analyzed with a gas chromatograph with a flame ionization detector and a fused silica column. Samples were periodically obtained for analysis by gas chromatography/ mass spectroscopy.

* * Sham-exposed and test animals were housed in their exposure chambers throughout the exposure period; untreated controls in a separate room. Food and water were not available during the 6 -hour exposure periods but were available ad libitum at all other times. All animals were observed daily. Body weights were recorded on days 0, 6, 13, 16, and 20. Food consumption was measured on GD 6, 13, 16, and 20. Females were sacrificed on GD 20 by diethyl ether overexposure followed by exsanguination. samples from the descending aorta were analyzed for glucose, urea nitrogen, total protein, albumin, globulin, A/G ratio, sorbitol dehydrogenase, lactate dehydrogenase, aspartate aminotransferase, alanine aminotransferase, alkaline phosphatase, total bilirubin, creatinine, cholesterol, triglycerides, uric acid, Cl, Ca, Na, K, and P. All organs were examined grossly. The number of corpora lutea per ovary and the gravid uterine weights were recorded. Uterine contents were examined and the numbers of implantation sites, early and late resorptions, and live and dead fetuses were recorded. The gender of each fetus was recorded. Fetuses were weighed and examined for gross anomalies. Fetuses of each litter were equally distributed between two groups; half were fixed in Bouin's solution and examined for visceral anomalies and the remaining fetuses were fixed in 95% ethanol and examined for skeletal anomalies after differential staining for cartilage and bone. * *

** Dams exposed to 6745 ppm had a slight reduction in body weight gain and a significant decrease in food consumption. A concentration-related increase in the incidence of rudimentary 14th ribs was observed (statistically significant at 3095 and 6745 ppm) but the relevance of the finding was uncertain. There was no apparent toxicity, either maternal or fetal, at the lowest exposure concentration, 430 ppm.

F020 241318 EOR F002 455 F010 5.8.2 F004 2 F005 RM

* *

* *

```
F006 Type: Developmental Toxicity
    Species/strain: Sprague-Dawley; VAF/Plus Crl:CD(SD)BR
* *
    No./dose: 22/group
* *
    Vehicle: None
* *
    Method: USEPA 1984; 40CFR Part 798:4350
F007 Type: Developmental Toxicity
    Species/strain: Sprague-Dawley; VAF/Plus Crl:CD(SD)BR
* *
    No./dose: 22/group
    Vehicle: None
* *
    Method: USEPA 1984; 40CFR Part 798:4350
F020 241322
EOR
F002 455
F010 5.8.2
F004 2
F005 RS
F006 Maternal NOEL: 430 ppm
** Pup NOEL: 430 ppm
F007 Maternal NOEL: 430 ppm
** Pup NOEL: 430 ppm
F020 241317
EOR
F002 455
F010 5.8.2
F004 2
F005 TS
F006 Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.
F007 Diisopropyl Ether (CAS No. 108-20-3); purity ~92%.
F020 241315
EOB
С
Χ
```

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C International Uniform Chemical Information Database
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C Company
           : ExxonMobil Biomedical Sciences Inc. 08801-3059 Annadale, New Je
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F003 Y27-002
F004 201-158-5
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F002 27
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F001 12032693
F003 ExxonMobil Biomedical Sciences Inc.
F004 1545 Route 22 East
F005 Annadale, New Jersey
F006 08801-3059
F008 A31-024
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F004 3
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F006 17-01-2002
F007 14-01-2002
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F004 2
F005 2
F006 17-01-2002
F007 14-01-2002
EOR
F001 27
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F005 6
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F007 17-01-2002
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F004 7
F005 7
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F007 17-01-2002
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F002 0
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F005 6
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F004 7
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F004 4
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F007 14-01-2002
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F003 4.3
F004 4
F005 4
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EOR
F001 27
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F006 17-01-2002
F007 14-01-2002
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F001 27
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F003 4.3

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F006 18-01-2002
F007 14-01-2002
EOR
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EOR
F001 27
F002 0
F003 5.0
F004 3
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F001 27
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F003 5.1.1
F004 4
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EOR
F001 27
F002 0
F003 5.1.1
F004 5
F005 5
F006 10-01-2002
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EOR
F001 27
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F003 5.1.2
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F005 2
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F003 5.1.2
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F003 5.1.3
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F005 1
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F005 1
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EOR
F001 27
F002 0
F003 5.2.1
F004 2
F005 2
F006 10-01-2002
F007 10-01-2002
EOR
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F003 5.2.2
F004 2
F005 2
F006 10-01-2002
F007 10-01-2002
EOR
F001 27
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F004 1
F005 1
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F007 11-01-2002
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F004 2
F005 2
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F007 11-01-2002
EOR
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F003 5.6
F004 1
F005 1
F006 11-01-2002
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F007 11-01-2002
EOR
F001 27
F002 0
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F004 2
F005 2
F006 29-01-2002
F007 29-01-2002
EOR
F001 27
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F003 78-92-2
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B115 GI_COMPANY_TAB
F001 27
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F008 Shell Nederland Chemie B.V.
F009 P.O. Box 3030
F010 Hoogvliet-Rotterdam
F011 3190 GH
F013 A31-015
F014 +31-10-2317005
F016 +31-10-2317125
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F001 27
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F008 Union Carbide Benelux
F009 Norderlaan 147
F010 Antwerpen
F011 2030
F013 A31-003
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F003 11-02-2000
F008 Atochem
F009 4, Cours Michelet
F010 Paris la Defense
F011 92080
F013 A31-007
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F009 89 bld Franklin Roosevelt
F010 Rueil Malmaison
F011 92564
F013 A31-007
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F015 SHELL 615013F
F016 33 1 47.14.82.99
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F009 4600 Parkway
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F013 A31-023
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F008 Deutsche Shell Chemie GmbH
F009 Koelner Strasse 6
F010 Eschborn
F011 65760
F013 A31-008
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F008 RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
F009 Ueberseering 40
F010 Hamburg
F011 22297
F013 A31-008
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F015 21151320
F016 040-6375-3496
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F011 A19-02
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B102 GI_SYNONYM_TAB
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F007 2-Butanol
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F004 HEDSET
F007 Ethyl methyl carbinol
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F003 19-05-1994
F004 HEDSET
F007 2-Hydroxybutane
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F007 sec-butanol
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F004 HEDSET
F007 2-butanol
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F001 27
F002 6
F003 26-05-1994
F004 HEDSET
F007 secundary butyl alcohol
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F003 27-04-1994
F004 HEDSET
F007 SBA; sec-Butanol; Butan-2-ol; secondary butyl alcohol
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F003 20-05-1994
F004 HEDSET
F007 SBA; secondary butyl alcohol; ethyl methyl carbinol; 2-hydroxybutane
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F004 DS_COMPARE_PKG
F007 SBA
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F007 butan-2-ol
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F004 DS_COMPARE_PKG
F007 secondary butyl alcohol
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F007 sec.-butyl alcohol, iso-butanol
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F016 A12-03
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B108 GI_CATEGORY_TAB
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F004 IUC300 COL
F007 A20-01
F008 A14-05
EOR
F001 27
F002 40
F003 11-02-2000
F004 IUC300_COL
F007 A20-01
F008 A14-08
EOR
F001 27
F002 41
F003 11-02-2000
F004 IUC300_COL
F007 A20-02
F008 A13-01
EOR
F001 27
F002 42
F003 11-02-2000
F004 IUC300 COL
F007 A20-02
F008 A13-02
EOR
F001 27
F002 43
F003 11-02-2000
F004 IUC300_COL
F007 A20-02
F008 A13-03
EOR
F001 27
F002 44
F003 11-02-2000
F004 IUC300_COL
F007 A20-02
F008 A13-04
EOR
F001 27
F002 45
F003 11-02-2000
F004 IUC300_COL
F007 A20-03
F008 A15-33
EOR
F001 27
F002 46
F003 11-02-2000
F004 IUC300 COL
F007 A20-03
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F008 A15-48

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EOR
F001 27
F002 47
F003 11-02-2000
F004 IUC300_COL
F007 A20-03
F008 A15-55
EOR
F001 27
F002 48
F003 11-02-2000
F004 IUC300_COL
F007 A20-03
F008 A15-55: fuel additive in lead-free gasoline
EOB
B109 GI_EXPO_LIMIT_TAB
F001 27
F002 2
F003 03-06-1994
F004 HEDSET
F007 A17-03
F008 450
F009 A16-03
EOR
F001 27
F002 3
F003 03-06-1994
F004 HEDSET
F007 A17-07
F008 303
F009 A16-03
EOR
F001 27
F002 4
F003 26-05-1994
F004 HEDSET
F007 A17-07
F008 303
F009 A16-03
F010 455
F011 A16-03
EOR
F001 27
F002 5
F003 27-04-1994
F004 HEDSET
F007 A17-07
F008 303
F009 A16-03
F010 455
F011 A16-03
F012 15
F013 A18-02
F014 4
EOR
F001 27
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F002 6
F003 27-04-1994
F004 HEDSET
F007 A17-04
F008 300
F009 A16-03
F010 600
F011 A16-03
F012 30
F013 A18-02
F014 4
EOR
F001 27
F002 7
F003 27-04-1994
F004 HEDSET
F007 A17-09: VME
F008 300
EOR
F001 27
F002 8
F003 26-05-1994
F004 HEDSET
F007 A17-07
F008 303
F009 A16-03
EOR
F001 27
F002 9
F003 16-02-1994
F004 DS_COMPARE_PKG
F007 A17-07
F008 303
F009 A16-03
F010 455
F011 A16-03
F012 8
F013 A18-01
EOR
F001 27
F002 10
F003 16-02-1994
F004 DS_COMPARE_PKG
F007 A17-04
F008 300
F009 A16-03
EOR
F001 27
F002 11
F003 16-02-1994
F004 HEDSET
F007 A17-03
F008 450
F009 A16-03
EOR
F001 27
```

F002 12

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F003 16-02-1994
F004 DS_COMPARE_PKG
F007 A17-06
F008 300
F009 A16-03
F010 450
F011 A16-03
EOR
F001 27
F002 13
F003 30-05-1994
F004 HEDSET
F007 A17-04
F008 100
F009 A16-04
EOR
F001 27
F002 16
F003 13-05-1994
F004 HEDSET
F007 A17-03
F008 450
F009 A16-03
EOB
B110 GI_SOURCE_OF_EXPOSURE_TAB
F001 27
F002 1
F003 24-05-1994
F004 HEDSET
EOR
F001 27
F002 2
F003 03-06-1994
F004 HEDSET
EOR
F001 27
F002 3
F003 23-02-1994
F004 HEDSET
EOB
B114 GI_OTHER_TAB
F001 27
F002 1
F003 10-06-1994
F004 HEDSET
EOR
F001 27
F002 2
F003 26-05-1994
F004 HEDSET
EOR
F001 27
F002 3
F003 26-05-1994
F004 HEDSET
```

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EOR
F001 27
F002 4
F003 30-05-1994
F004 HEDSET
EOB
B201 PC_MELTING_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F016 2
F007 A02-06
F008 -89
F009 -108
F012 P01-03: no data
F014 A03-02
EOR
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F016 1
F007 A02-03
F008 -114
F012 P01-03: no data
F014 A03-02
EOB
B202 PC_BOILING_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F017 2
F007 A02-06
F008 99
F009 102
F010 1013
F011 P02-01
F013 P03-03: ASTM D1078/86
F014 1986
F015 A03-02
EOR
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F017 1
F007 A02-03
F008 99.5
F013 P03-03: no data
F015 A03-02
EOB
B203 PC_DENSITY_TAB
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F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F017 2
F007 P05-02
F008 A02-06
F009 807
F011 P18-02
F012 15
F013 P04-03: ASTM D4052/86
F014 1986
F015 A03-02
EOR
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F007 P05-02
F008 A02-03
F009 3.29
F011 P18-02
F012 20
F013 P04-03: no data
F015 A03-02
EOR
F001 27
F002 3
F003 17-01-2002
F004 DAWINKE
F017 1
F007 P05-02
F008 A02-03
F009 806
F011 P18-02
F012 20
F013 P04-03: no data
F015 A03-02
EOB
B204 PC_VAPOUR_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F016 1
F007 A02-03
F008 15.98
F010 P02-01
F011 20
F012 P06-04
F014 A03-02
EOB
B205 PC_PARTITION_TAB
F001 27
F002 2
```

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F003 17-01-2002
F004 DAWINKE
F015 1
F007 A02-03
F008 .61
F010 20
F011 P07-05
F013 A03-02
EOB
B206 PC_WATER_SOL_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F024 3
F007 A02-06
F008 P08-01
F009 201
F011 20
F020 P09-03: no data
F022 A03-02
EOR
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F024 1
F007 A02-03
F008 P08-01
F009 125
F011 20
F020 P09-03: no data
F022 A03-02
EOR
F001 27
F002 3
F003 17-01-2002
F004 DAWINKE
F024 2
F007 A02-03
F008 P08-01
F009 181
F011 25
F020 P09-03: no data
F022 A03-02
EOB
B207 PC_FLASH_TAB
F001 27
F002 1
F003 01-06-1994
F004 HEDSET
F007 A02-06
F008 25
F009 P10-01
F010 P11-01
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F011 1987
F012 A03-02
EOR
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F014 1
F007 A02-03
F008 24
F009 P10-01
F010 P11-02: NFT 60 103
F012 A03-02
EOB
B208 PC_AUTO_FLAMM_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F016 1
F007 A02-04
F008 350
F010 1013
F011 P02-01
F012 P13-03: E695/85
F013 1985
F014 A03-02
EOR
F001 27
F002 2
F003 02-06-1994
F004 HEDSET
F007 A02-03
F008 406
F010 1013
F011 P02-01
F012 P13-03: no data
F014 A03-02
EOB
B209 PC_FLAMM_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F012 1
F007 P16-04
F008 P15-05: no data
F010 A03-02
EOB
B210 PC_EXPL_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
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F012 1
F008 P21-02: no data
F010 A03-02
EOB
C
B301 EN PHOTODEGRADATION TAB
F001 27
F002 1
F003 18-01-2002
F004 DAWINKE
F046 5
EOR
F001 27
F002 4
F003 17-01-2002
F004 DAWINKE
F046 4
F008 F01-01
F009 F02-06
F011 F03-03: sun lamp
F043 A03-02
EOR
F001 27
F002 5
F003 17-01-2002
F004 DAWINKE
F046 3
F008 F01-01
F009 F02-04
F010 1990
F034 F06-03
F035 1000000
F036 F07-02
F044 A02-03
F037 .000000000096466
F038 A02-03
F040 50
F041 20
F042 F05-02
EOR
F001 27
F002 6
F003 17-01-2002
F004 DAWINKE
F045 A36-003
F046 1
F008 F01-01
F009 F02-05
F034 F06-03
F035 1500000
F036 F07-02
F044 A02-03
F037 .000000000099751
F038 A02-03
F040 50
F041 12.9
F042 F05-02
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EOR
F001 27
F002 7
F003 17-01-2002
F004 DAWINKE
F046 2
F008 F01-01
F009 F02-06
F034 F06-03
F043 A03-02
EOB
С
B302 EN_STABILITY_IN_WATER_TAB
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F041 1
F008 F08-01
F039 A03-02
EOB
B303 EN_STABILITY_IN_SOIL_TAB
F001 27
F002 1
F003 18-01-2002
F004 DAWINKE
F055 1
EOB
B304 EN_MONITORING_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F010 6
F007 F19-01
F008 F35-09
EOR
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F010 1
F007 F19-01
F008 F35-07
EOR
F001 27
F002 3
F003 17-01-2002
F004 DAWINKE
F010 4
F007 F19-02
F008 F35-08: wastewater
EOR
F001 27
F002 4
```

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F003 17-01-2002
F004 DAWINKE
F010 5
F007 F19-02
F008 F35-08: landfill leachate
EOR
F001 27
F002 5
F003 17-01-2002
F004 DAWINKE
F010 2
F007 F19-01
F008 F35-01
EOR
F001 27
F002 6
F003 17-01-2002
F004 DAWINKE
F010 3
F007 F19-01
F008 F35-01
EOB
B305 EN_TRANSPORT_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F012 1
F007 F20-03
F008 F22-03
F009 F21-01
EOB
B306 EN_DISTRIBUTION_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F010 A36-003
F011 1
F007 F24-02
F008 F23-01
EOB
B308 EN_BIODEGRADATION_TAB
F001 27
F002 2
F003 14-01-2002
F004 CLGETTS
F048 7
F007 A01-01
F008 F25-01
F011 F27-0166: semi-automatic activated sludge system
F020 F30-02: 98% BOD reduction after 5 days
F046 A03-02
EOR
```

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F001 27
F002 3
F003 14-01-2002
F004 CLGETTS
F048 9
F007 A01-01
F008 F25-02
F011 F27-0166: acetate-acclimated methane cultures
F017 100
F018 14
F019 F05-01
F046 A03-02
EOR
F001 27
F002 4
F003 14-01-2002
F004 CLGETTS
F048 8
F007 A01-01
F008 F25-02
F011 F27-0166: acetate enriched cultures
F046 A03-02
EOR
F001 27
F002 5
F003 17-01-2002
F004 DAWINKE
F048 5
F008 F25-01
F009 F26-25: screening test
F011 F27-0137
F015 A02-03
F017 33
F018 5
F019 F05-01
F046 A03-02
EOR
F001 27
F002 7
F003 17-01-2002
F004 DAWINKE
F048 4
F008 F25-01
F009 F26-25: screening test
F011 F27-0137
F015 A02-03
F017 81.7
F018 5
F019 F05-01
F046 A03-02
EOR
F001 27
F002 8
F003 17-01-2002
F004 DAWINKE
F048 6
F008 F25-01
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F009 F26-25: screening test
F011 F27-0137
F015 A02-03
F017 9.3
F018 1
F019 F05-01
F046 A03-02
EOR
F001 27
F002 9
F003 17-01-2002
F004 DAWINKE
F048 3
F008 F25-01
F009 F26-25: screening test
F011 F27-0137
F015 A02-03
F017 98.5
F018 5
F019 F05-01
F046 A03-02
EOR
F001 27
F002 10
F003 18-01-2002
F004 DAWINKE
F047 A36-003
F048 1
F008 F25-01
F009 F26-25: APHA (American Public Health Association) Standard Methods, No.
     219
F011 F27-0166: Wastewater treatment plant
F015 A02-03
F017 83
F018 5
F019 F05-01
F046 A03-02
F052 5
F053 F05-01
EOR
F001 27
F002 11
F003 18-01-2002
F004 DAWINKE
F047 A36-003
F048 2
F009 F26-25: Winkler Method
F011 F27-0166: Wastewater treatment plant
F046 A03-02
F052 50
F053 F05-01
EOB
B309 EN BOD COD TAB
F001 27
F002 1
F003 17-01-2002
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F004 DAWINKE
F023 1
F007 F32-03: not specified
F009 A03-02
F013 A02-03
F014 1.87
F015 F33-03: not specified
F017 A03-02
F018 A02-03
F019 2.47
F020 A02-03
F021 .76
EOB
B310 EN_BIOACCUMULATION_TAB
F001 27
F002 1
F003 18-01-2002
F004 DAWINKE
F022 1
F009 F34-06: calculated value
F016 A02-06
F017 1.71
F020 A03-02
EOB
B311 EN_OTHER_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F010 1
F009 The volatilization half-life in a model river is estimated to be 3.5 days
     at 25 degrees C.
EOB
B401 EC FISHTOX TAB
F001 27
F002 3
F003 17-01-2002
F004 DAWINKE
F034 3
F008 E01-05
F009 E02-0075
F010 E03-05: not specified
F012 48
F013 E04-02
F014 E05-02
F021 A02-03
F022 3520
F031 A03-01
F032 A03-02
F045 E35-02
EOR
F001 27
F002 4
F003 18-01-2002
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F004 DAWINKE
F034 4
F008 E01-05
F009 E02-0113
F010 E03-05: not specified
F012 24
F013 E04-02
F031 A03-01
F032 A03-02
EOR
F001 27
F002 5
F003 18-01-2002
F004 DAWINKE
F033 A36-003
F034 1
F007 A01-01
F008 E01-02
F009 E02-0119
F010 E03-05: no data
F011 1985
F012 96
F013 E04-02
F014 E05-02
F021 A02-03
F022 3670
F031 A03-03
F032 A03-02
F045 E35-02
EOR
F001 27
F002 6
F003 17-01-2002
F004 DAWINKE
F033 A36-003
F034 2
F008 E01-05
F009 E02-0015
F010 E03-05: APHA (American Public Health Association) Standard Methods
F012 24
F013 E04-02
F014 E05-02
F021 A02-03
F022 4300
F031 A03-03
F032 A03-02
F045 E35-02
EOR
F001 27
F002 7
F003 17-01-2002
F004 DAWINKE
F033 A36-003
F034 5
F009 E02-0161: fish
F010 E03-05: ECOSAR Computer Model
F012 96
```

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F013 E04-02
F014 E05-02
F021 A02-03
F022 1113
F045 E35-01
EOB
B402 EC_DAPHNIATOX_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F033 2
F008 E06-0010
F009 E07-04: not specified
F011 24
F012 E04-02
F013 E05-02
F026 LC50
F027 A02-03
F028 3750
F030 A03-01
F031 A03-02
F042 E01-05
F047 E35-02
EOR
F001 27
F002 2
F003 18-01-2002
F004 DAWINKE
F032 A36-003
F033 1
F007 A01-01
F008 E06-0010
F009 E07-04: German Institute of Standardization, DIN 38412, Part II, Daphnia
     Short-Time Test
F011 48
F012 E04-02
F013 E05-02
F020 A02-03
F021 4227
F030 A03-01
F031 A03-02
F042 E01-05
F045 E35-02
EOR
F001 27
F002 3
F003 17-01-2002
F004 DAWINKE
F032 A36-003
F033 3
F008 E06-0034: Tetrahymena pyrifomis (Ciliate)
F009 E07-04: Population Growth Impairment Test
F011 48
F012 E04-02
F013 E05-02
```

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F020 A02-03
F021 3196
F030 A03-02
F031 A03-02
F042 E01-05
F045 E35-02
EOR
F001 27
F002 4
F003 17-01-2002
F004 DAWINKE
F032 A36-003
F033 4
F008 E06-0034: Daphnid
F009 E07-04: ECOSAR Computer Model
F011 48
F012 E04-02
F013 E05-02
F026 LC50
F027 A02-03
F028 1084
F047 E35-01
EOB
B403 EC_ALGAETOX_TAB
F001 27
F002 2
F003 18-01-2002
F004 DAWINKE
F037 4
F008 E08-0037
F009 E09-04: not specified
F012 192
F013 E04-02
F014 E05-02
F015 A02-03
F016 312
F034 A03-02
F035 A03-02
F046 E35-02
EOR
F001 27
F002 3
F003 18-01-2002
F004 DAWINKE
F037 5
F008 E08-0018
F009 E09-04: not specified
F014 E05-02
F015 A02-03
F016 8900
F034 A03-02
F035 A03-02
F046 E35-02
EOR
F001 27
F002 4
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F003 17-01-2002
F004 DAWINKE
F036 A36-003
F037 1
F008 E08-0053
F009 E09-04: 7-Day Cell Multiplication Inhibition Test
F012 7
F013 E04-01
F014 E05-02
F030 TT
F031 A02-03
F032 95
F034 A03-01
F035 A03-02
F051 E35-02
EOR
F001 27
F002 5
F003 17-01-2002
F004 DAWINKE
F036 A36-003
F037 2
F008 E08-0063: green alga
F009 E09-04: ECOSAR Computer Model
F012 96
F013 E04-02
F014 E05-02
F027 A02-03
F028 625
F050 E35-01
EOR
F001 27
F002 6
F003 18-01-2002
F004 DAWINKE
F036 A36-003
F037 3
F009 E09-04: ECOSAR Computer Model
F011 E10-03: green alga
F012 96
F013 E04-02
F014 E05-02
F030 ChV*
F031 A02-03
F032 28
F051 E35-01
EOB
B404 EC_BACTOX_TAB
F001 27
F002 1
F003 18-01-2002
F004 DAWINKE
F035 1
F008 E29-01
F009 E11-0109
F010 E12-06: total biomass
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F012 16
F013 E04-02
F014 E05-02
F015 A02-06
F016 500
F024 NOAEL
F025 A02-03
F026 500
F032 A03-02
F033 A03-02
F036 E35-02
EOR
F001 27
F002 2
F003 18-01-2002
F004 DAWINKE
F035 2
F009 E11-0029
F014 E05-02
F021 A02-03
F022 1630
F032 A03-02
F033 A03-02
F038 E35-02
EOR
F001 27
F002 3
F003 17-01-2002
F004 DAWINKE
F035 5
F008 E29-01
F009 E11-0140
F010 E12-06: total biomass
F012 20
F013 E04-02
F014 E05-02
F024 NOEC
F025 A02-03
F026 1416
F032 A03-02
F033 A03-02
EOR
F001 27
F002 4
F003 18-01-2002
F004 DAWINKE
F035 3
F008 E29-01
F009 E11-0037
F010 E12-06: total biomass
F012 48
F013 E04-02
F014 E05-02
F024 NOEL
F025 A02-03
F026 745
F032 A03-02
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F033 A03-02
F039 E35-02
EOR
F001 27
F002 5
F003 18-01-2002
F004 DAWINKE
F035 4
F008 E29-01
F009 E11-0054
F010 E12-06: total biomass
F012 72
F013 E04-02
F014 E05-02
F024 NOEL
F025 A02-03
F026 1282
F032 A03-02
F033 A03-02
F039 E35-02
EOB
B405 EC_CHRONFISHTOX_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F030 A36-003
F031 1
F008 E02-0161: fish
F009 E13-02: ECOSAR Computer Model
F012 30
F013 E15-01
F014 E05-02
F024 ChV*
F025 A02-03
F026 115
F035 E35-01
EOB
B406 EC_CHRONDAPHNIATOX_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F030 A36-003
F031 1
F008 E06-0034: Daphnid
F009 E16-02: ECOSAR Computer Model
F012 16
F013 E18-01
F014 E05-02
F015 A02-03
F016 30
F032 E35-01
EOB
C
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B408 EC_PLANTTOX_TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F034 5
F008 E23-06
F009 E24-02: not indicated
F011 E25-03: seed germination
F014 E05-02
F018 A02-03
F019 650
F032 A03-02
EOR
F001 27
F002 2
F003 17-01-2002
F004 DAWINKE
F034 6
F008 E23-17: Cucumis sativus
F009 E24-02: not specified
F011 E25-03: seed germination
F014 E05-02
F015 A02-01
F016 50375
F032 A03-02
EOR
F001 27
F002 3
F003 17-01-2002
F004 DAWINKE
F034 4
F008 E23-17: Lupinus albus
F012 1
F013 E15-01
F032 A03-02
EOR
F001 27
F002 4
F003 17-01-2002
F004 DAWINKE
F034 1
F008 E23-17: Solanum tuberosum L.
F032 A03-02
EOR
F001 27
F002 5
F003 17-01-2002
F004 DAWINKE
F034 3
F008 E23-17: wheat
F032 A03-02
EOR
F001 27
F002 6
F003 17-01-2002
F004 DAWINKE
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F034 2
F008 E23-17: wheat
F032 A03-02
EOB
C
B407 EC SOILDWELLINGTOX TAB
F001 27
F002 1
F003 17-01-2002
F004 DAWINKE
F033 A36-003
F034 1
F009 E20-0039: earthworm
F010 E21-03: ECOSAR Computer Model
F012 E22-01
F013 14
F014 E04-01
F015 E33-03: ppm
F022 A02-03
F023 1222
F037 E35-01
EOB
B409 EC_OTHER_SPEC_TAB
F001 27
F002 2
F003 18-01-2002
F004 DAWINKE
F032 A36-003
F008 E26-15: Xenopus laevis (Clawed Frog)
F009 E27-03: not specified
F011 E28-01
F012 48
F013 E30-03
F014 E32-02:
             mg/L
F021 A02-03
F022 1530
F031 A03-02
F036 E35-02
EOB
B017 TO_META_MAM_TAB
F001 27
F002 1
F003 01-06-2006
F004 CLGETTS
F005 A36-002
F006 1
F009 T02-24
F021 T52-003: Aqueous Solution
F022 C34-001: Oral Gavage / i.v.
F034 C36-001: Experimental Model Development
F035 1981
F036 A03-02
F037 A01-03: 2-Butanol (sec-Butanol; sBA) and metabolites 2-Butanone (Methyl
     ethyl ketone; MEK), 3-Hydroxy-2-Butanone (3H-2B), and 2,3-Butanediol
     (2,3-BD) (CAS Nos. 78-92-2 (sBA); 78-93-3 (MEK))
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F042 (See Remark)
EOR
F001 27
F002 2
F003 01-06-2006
F004 CLGETTS
F005 A36-002
F006 2
F009 T02-10
F021 T52-003: Corn oil (25% solution)
F022 C34-003
F034 C36-001: Experimental
F035 1976
F036 A03-02
F037 A01-03: 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)
F042 450 mg/kg body weight
EOR
F001 27
F002 3
F003 01-06-2006
F004 CLGETTS
F005 A36-002
F006 3
F009 T02-13
F010 9
F021 T52-003: none
F022 C34-010
F034 C36-001: Experimental
F035 1988
F036 A03-02
F037 A01-03: 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)
F042 200 ppm (8.2mmol/m3)
EOB
С
B501 TO_ACUTE_ORAL_TAB
F001 27
F002 1
F003 21-10-1992
F004 HEDSET
F007 A01-01
F008 T01-03
F009 T02-24
F010 T03-03
F012 A02-03
F013 6500
F015 T04-01
F016 A03-02
EOR
F001 27
F002 2
F003 21-10-1992
F004 HEDSET
F007 A01-01
F008 T01-03
F009 T02-23
F010 T03-03: not specified
F011 1972
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F012 A02-03
F013 4900
F015 T04-01
F016 A03-02
EOR
F001 27
F002 3
F003 10-01-2002
F004 CLGETTS
F017 A36-002
F007 A01-01
F008 T01-03
F009 T02-24
F010 T03-03: Experimental (Non-regulatory)
F011 1986
F012 A02-03
F013 2054
F014 2328
F015 T04-01
F016 A03-03
F019 T24-03
F020 10
F021 T52-003: None
F022 T23-16
EOR
F001 27
F002 4
F003 10-01-2002
F004 CLGETTS
F017 A36-003
F007 A01-01
F008 T01-03
F009 T02-24
F010 T03-03: Experimental (Non-regulatory)
F011 1954
F012 A02-03
F013 5730
F014 7320
F015 T04-01
F016 A03-01
F019 T24-02
F020 5
F021 T52-003: Unknown (water, corn oil, or a 1% solution of
     3,9-diethyl-6-tridecanol sulfate (Tergitol Penetrant 7)
F022 T23-48: Carworth-Wistar
EOR
F001 27
F002 5
F003 10-01-2002
F004 CLGETTS
F017 A36-004
F007 A01-01
F008 T01-03
F009 T02-23
F010 T03-03: Experimental (Non-regulatory)
F011 1925
F012 A02-03
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F013 4890
F015 T04-01
F016 A03-01
F019 T24-03
F021 T52-002
F022 T23-48: Unknown
EOB
C
B502 TO_ACUTE_INHAL_TAB
F001 27
F002 1
F003 11-01-2002
F004 CLGETTS
F019 A36-003
F007 A01-01
F008 T05-03
F009 T02-24
F010 T06-03: Experimental (Non-regulatory)
F011 1954
F012 A02-03
F013 16000
F015 T07-02
F016 4
F017 T08-01
F018 A03-01
F021 T24-02
F022 6
F023 T52-003: None
F024 T23-48: Carworth-Wistar
EOR
F001 27
F002 2
F003 10-01-2002
F004 CLGETTS
F019 A36-003
F007 A01-01
F008 T05-03
F009 T02-24
F010 T06-03: Experimental (Non-regulatory)
F011 1951
F012 A02-03
F013 8000
F015 T07-02
F016 4
F017 T08-01
F018 A03-01
F021 T24-02
F022 6
F023 T52-003: None
F024 T23-48: Carworth-Wistar
EOR
F001 27
F002 3
F003 10-01-2002
F004 CLGETTS
F019 A36-003
F007 A01-01
```

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F008 T05-03
F009 T02-24
F010 T06-03: Experimental (Non-regulatory)
F011 1989
F012 A02-03
F013 10000
F015 T07-02
F016 7
F017 T08-01
F018 A03-03
F021 T24-01
F022 5
F023 T52-003: None
F024 T23-42
EOR
F001 27
F002 4
F003 10-01-2002
F004 CLGETTS
F019 A36-005
F007 A01-01
F008 T05-03
F009 T02-18
F010 T06-03: Experimental (Non-regulatory)
F011 1938
F018 A03-01
F021 T24-04
F022 2
F023 T52-003: None
F024 T23-47
EOB
С
B503 TO_ACUTE_DERMAL_TAB
F001 27
F002 1
F003 10-01-2002
F004 CLGETTS
F017 A36-002
F007 A01-01
F008 T01-05: Limit
F009 T02-24
F010 T09-02: Experimental (Non-regulatory)
F011 1986
F012 A02-04
F013 2000
F015 T04-01
F016 A03-03
F019 T24-03
F020 10
F021 T52-003: None
F022 T23-16
EOB
B504 TO ACUTE OTHER TAB
F001 27
F002 1
F003 21-10-1992
```

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F004 HEDSET
F007 A01-01
F008 T10-07
F009 T02-18
F010 T11-02
F011 other: not specified
F012 1978
F013 A02-03
F014 800
F016 T12-01
F019 A03-02
EOB
B505 TO_SKIN_IRRITATION_TAB
F001 27
F002 1
F003 21-10-1992
F004 HEDSET
F007 A01-01
F008 T02-23
F009 T14-06: not specified
F010 1954
F011 T15-04
F012 T46-06
F013 A03-02
EOR
F001 27
F002 2
F003 10-01-2002
F004 CLGETTS
F014 A36-002
F007 A01-01
F008 T02-23
F009 T14-05
F010 1986
F011 T15-04
F012 T46-06
F013 A03-03
F018 T50-001
F019 4
F020 T55-001
F021 6
EOB
B506 TO_EYE_IRRITATION_TAB
F001 27
F002 1
F003 10-01-2002
F004 CLGETTS
F014 A36-003
F007 A01-01
F008 T02-23
F009 T16-04:
             Experimental (Non-regulatory)
F010 1954
F011 T17-01
F012 T46-04
F013 A03-01
```

```
F022 5
EOR
F001 27
F002 2
F003 10-01-2002
F004 CLGETTS
F014 A36-002
F007 A01-01
F008 T02-23
F009 T16-03
F010 1986
F012 T46-01
F013 A03-03
F022 6
EOB
B507 TO_SENSITIZATION_TAB
F001 27
F002 1
F003 11-01-2002
F004 CLGETTS
F015 A36-002
F007 A01-01
F008 T18-03
F009 T02-10
F010 T20-03: Magnusson and Kligman; 1969
F011 1986
F012 T47-01
F013 T21-02
F014 A03-03
F017 20
F030 T52-003: Corn oil
EOR
F001 27
F002 2
F003 11-01-2002
F004 CLGETTS
F015 A36-002
F007 A01-03
F008 T18-04
F009 T02-10
F010 T20-03: OECD Guide-line 406 and EC92/69/EEC, B6
F011 1992
F012 T47-01
F013 T21-02
F014 A03-03
F017 20
F030 T52-003: Paraffin oil
EOB
B508 TO_REPEATED_DOSE_TAB
F001 27
F002 1
F003 14-01-2002
F004 CLGETTS
F030 A36-002
F031 2
```

```
F007 A01-03
F008 T02-24
F009 T23-16
F010 T24-03
F011 T25-08
F012 T26-16:
             Comparable to OECD Guideline 413; 90-Day Subchronic Toxicity
F013 1981
F014 90 days
F015 6 hours/day; 5 days/week
F017 0, 1250, 2500, or 5000 ppm - Vapor
F018 T27-07
F019 A02-03
F020 2500
F022 T28-05
F024 A02-03
F025 5000
F027 T28-05
F029 A03-03
EOR
F001 27
F002 2
F003 11-01-2002
F004 CLGETTS
F030 A36-003
F031 1
F007 A01-01
F008 T02-24
F009 T23-42
F010 T24-02
F011 T25-08
F012 T26-16: Experimental -- Study of changes in cytochrome P-450 enzyme
     system of rat liver, kidney and lung
F013 1985
F014 3-5 days
F015 6 hours/day
F017 2000 ppm (3 Days) and 500 ppm (5 Days)
F029 A03-02
EOB
B509 TO_GENETIC_IN_VITRO_TAB
F001 27
F002 1
F003 11-01-2002
F004 CLGETTS
F016 A36-002
F007 A01-01
F008 T30-01
F009 T31-10
F010 1985
F011 Bacterial
F012 T32-03
F013 T33-02
F014 A03-03
F015 0.01, 0.1, 0.5, 1.0, and 5.0 mg/ml
F001 27
F002 2
```

```
F003 14-01-2002
F004 CLGETTS
F016 A36-002
F007 A01-01
F008 T30-01
F009 T31-11
F010 1985
F011 Bacterial
F012 T32-03
F013 T33-02
F014 A03-03
F015 0.01, 0.1, 0.5, 1.0, and 5.0 mg/ml
EOR
F001 27
F002 3
F003 14-01-2002
F004 CLGETTS
F016 A36-002
F007 A01-01
F008 T30-19: Yeast mitotic gene conversion
F009 T31-16
F010 1988
F011 Saccharomyces cerevisiae
F012 T32-03
F013 T33-02
F014 A03-03
F015 Maximum conc. 5 mg/ml
EOR
F001 27
F002 4
F003 14-01-2002
F004 CLGETTS
F016 A36-002
F007 A01-01
F008 T30-19: chromosome abberration assay
F009 T31-12
F010 1988
F011 Chinese hamster ovary cells
F012 T32-03
F013 T33-02
F014 A03-03
F015 maximum conc. 5000 ug/ml
EOR
F001 27
F002 5
F003 11-01-2002
F004 CLGETTS
F016 A36-002
F007 A01-01
F008 T30-01
F009 T31-10
F010 1988
F011 Bacterial
F012 T32-03
F013 T33-02
F014 A03-03
F015 100-500-1000-5000-10000 ug/plate
```

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EOB
B510 TO_GENETIC_IN_VIVO_TAB
F001 27
F002 1
F003 11-01-2002
F004 CLGETTS
F018 A36-005
F007 A01-01
F008 T34-12: Rat Bone Marrow
F009 T02-24
F010 T23-47
F011 T37-15: Micronucleus Aberration
F012 1988
F013 T24-04
F014 T25-11: intragastric
F017 A03-02
EOR
F001 27
F002 2
F003 29-01-2002
F004 CLGETTS
F018 A36-002
F007 A01-03
F008 T34-12: In Vivo Mammalian Bone Marrow Erythrocyte Micronucleus Assay
     Intraperitoneal Dosing Method
F009 T02-18
F010 T23-10
F011 T37-15: OECD 474 equivalent
F012 1988
F013 T24-03
F014 T25-11: Intraperitoneal injection
F015 12, 24 and 48 hours
F016 1.96 ml/kg Single injection
F017 A03-02
EOB
B511 TO_CARCINOGENICITY_TAB
F001 27
F002 1
F003 21-10-1992
F004 HEDSET
EOB
B512 TO_REPRODUCTION_TAB
F001 27
F002 1
F003 11-01-2002
F004 CLGETTS
F037 A36-003
F007 A01-01
F008 T41-04: two generation study with teratology screen
F009 T02-24
F010 T23-46
F011 T24-03
F012 T25-02
F013 T40-05: Comparable to an OECD 416 guideline study.
```

```
F014 1975
F015 Daily - ad libitum
F016 8 weeks
F017 8 weeks
F018 2 Generations
F019 F0 Generation: 0, 0.3, 1.0, or 3.0% solutions; F1 Generation: 0, 0.3,
    1.0, or 2.0%
F020 T27-07
F025 A02-03
F026 1
F028 T43-01
F035 A03-01
F039 T57-10: NOAEL Maternal
F040 A02-03
F041 1
F043 T43-01
EOB
B513 TO_DEVELOPMENTAL_TAB
F001 27
F002 1
F003 14-01-2002
F004 CLGETTS
F030 A36-003
F007 A01-01
F008 T02-24
F009 T23-42
F010 T24-01
F011 T25-08
F012 T44-03: Comparable to an OECD 414 guideline study
F013 1989
F014 20 days
F015 Gestation Days 1-19
F016 7 hr/day
F017 3500, 5000, 7000 ppm
F018 T27-07
F019 A02-03
F020 3500
F022 T43-04
F023 A02-04
F024 7000
F026 T43-04
F029 A03-03
EOR
F001 27
F002 2
F003 11-01-2002
F004 CLGETTS
F030 A36-003
F007 A01-01
F008 T02-24
F009 T23-46
F010 T24-01
F011 T25-02
F012 T44-03: Comparable to an OECD 421 quideline study.
F013 1975
F014 20 days
```

```
F015 8 Weeks premating (Males and Females) and during gestation (Females)
F016 Daily - ad libitum F017 0, 0.3, 1.0, or 2.0% solutions
F018 T27-07
F029 A03-01
EOB
B515 TO_HUMAN_EXPERIENCE_TAB
F001 27
F002 1
F003 22-10-1992
F004 HEDSET
EOB
B514 TO_OTHER_TAB
F001 27
F002 1
F003 11-01-2002
F004 CLGETTS
F008 A36-002
F007 T45-12: Respiratory Irritation
EOB
B601 TEXT_TAB
F002 27
F010 1.0.1
F004 7
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119787
EOR
F002 27
F010 1.0.1
F004 8
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119788
EOR
F002 27
F010 1.0.1
F004 9
F005 S0
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119789
EOR
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F002 27
F010 1.0.1
F004 10
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119790
EOR
F002 27
F010 1.0.1
F004 11
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119791
EOR
F002 27
F010 1.0.1
F004 12
F005 S0
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119792
EOR
F002 27
F010 1.0.1
F004 13
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119793
EOR
F002 27
F010 1.1.1
F004 8
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119794
EOR
F002 27
F010 1.10
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```
F004 1
F005 RM
F006 As the quantities of this substance placed on the EU market
     by Union Carbide Benelux N.V. are normally sourced from the
     manufacturing facilities of its U.S. parent company, no
* *
     exposure can arise within the EU from the manufacture of
* *
     these s
F007 As the quantities of this substance placed on the EU market
* *
     by Union Carbide Benelux N.V. are normally sourced from the
* *
     manufacturing facilities of its U.S. parent company, no
* *
     exposure can arise within the EU from the manufacture of
     these substances. The comments below on exposure are
* *
     restricted to uses for which Union Carbide believes its
* *
     customers use this substance.
* *
     Major use(s): Fuel additive in lead-free gasoline.
* *
* *
     Sources of human exposure: Very minor sporadic exposure to
* *
     the public via inhalation during filling of vehicles.
* *
     Quantitative estimates are not available.
* *
     Sources of environmental exposure: Negligible sporadic
* *
     exposure to the atmosphere during filling of vehicles.
* *
     Quantitative estimates are not available. Substance is
* *
     essentially oxidised to carbon dioxide and water during
* *
     use.
F008 HEDSET
F009 24-05-1994
F012 20
F020 119856
EOR
F002 27
F010 1.10
F004 1
F005 SO
F006 Union Carbide Benelux Antwerpen
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Union Carbide Benelux Antwerpen
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119857
EOR
F002 27
F010 1.10
F004 2
F006 Continuous process. Non-direct hydratation by absorption of
     butenes in sulfuric acid.
* *
     One production site.
* *
     Distribution pattern : from production
* *
                    0.02
         % to air
* *
         % to water 0.02
* *
         to soil
                     0.0
         to sediment 0.0
F007 Continuous process. Non-direct hydratation by absorption of
```

```
butenes in sulfuric acid.
* *
     One production site.
* *
     Distribution pattern : from production
* *
         % to air
                    0.02
* *
         % to water 0.02
* *
        to soil
                  0.0
* *
        to sediment 0.0
F008 HEDSET
F009 03-06-1994
F012 20
F020 119858
EOR
F002 27
F010 1.10
F004 2
F005 SO
F006 Atochem Paris la Defense
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Atochem Paris la Defense
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119859
EOR
F002 27
F010 1.10
F004 3
F005 RM
F006 Inhalation or skin contact when loading, unloading, using
     the product.
     In case of accidental release, product may contaminate the
* *
     environment.
F007 Inhalation or skin contact when loading, unloading, using
     the product.
* *
     In case of accidental release, product may contaminate the
* *
    environment.
F008 HEDSET
F009 23-02-1994
F012 20
F020 119860
EOR
F002 27
F010 1.10
F004 3
F005 SO
F006 SHELL FRANCE Rueil Malmaison
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 SHELL FRANCE Rueil Malmaison
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119861
EOR
F002 27
F010 1.11
```

```
F004 1
F005 RM
F006 DISPOSAL
* *
     Recover or recycle if possible. Otherwise incineration.
* *
* *
* *
     TRANSPORT INFORMATION
* *
* *
     UN Number: 1120
* *
     Class: 3
* *
     Packing Group: III
* *
     Proper Shipping Name: Secondary butyl alcohol
* *
* *
     Sea (IMO)
* *
     Class: 3.3
* *
     Packing Group: III
* *
     Symbol: Flammable
F007 DISPOSAL
* *
* *
     Recover or recycle if possible. Otherwise incineration.
* *
* *
* *
     TRANSPORT INFORMATION
* *
* *
     UN Number: 1120
* *
     Class: 3
* *
     Packing Group: III
     Proper Shipping Name: Secondary butyl alcohol
* *
     Sea (IMO)
* *
     Class: 3.3
* *
     Packing Group: III
* *
     Symbol: Flammable liquid
* *
     Marine Pollutant (Y/N): No
* *
* *
     Rail/Road (RID/ADR)
* *
     Class: 3
* *
     Item: 31(c)
* *
     Symbol: Flammable liquid
* *
     Kemler Plate: 30/1120
* *
* *
   Air (IATA/ICAO)
* *
     Class: 3
* *
     Packing Group: III
* *
     Symbol: Flammable liquid
F008 HEDSET
F009 03-06-1994
F012 20
F020 119795
EOR
F002 27
F010 1.11
F004 1
F005 SO
F006 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
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F007 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119796
EOR
F002 27
F010 1.11
F004 2
F005 RM
F006 Disposal: Incinerate in a furnace where permitted under
* *
     national and local regulations.
* *
* *
     Transport: Butan-2-ol is classified as a class 3 product
* *
     according the ADR/RID/IMDG/ICAO regulations.
* *
     Butan-2-ol is shipped in road/rail tankcars,
* *
F007 Disposal: Incinerate in a furnace where permitted under
* *
     national and local regulations.
* *
* *
     Transport: Butan-2-ol is classified as a class 3 product
* *
     according the ADR/RID/IMDG/ICAO regulations.
* *
     Butan-2-ol is shipped in road/rail tankcars,
* *
    tankcontainers/ISOtanks and smaller packages (e.g. drums).
* *
F008 HEDSET
F009 26-05-1994
F012 20
F020 119797
EOR
F002 27
F010 1.11
F004 2
F005 SO
F006 Union Carbide Benelux Antwerpen
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Union Carbide Benelux Antwerpen
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119798
EOR
F002 27
F010 1.11
F004 3
F006 Recover or recycle if possible. Otherwise : incineration.
     Avoid electrostatic discharge generation.
* *
     Earth all equipment.
* *
    Avoid splash filling.
* *
     Use a vapour recovery system.
* *
     Keep in a well ventilated place.
* *
     Avoid naked flames. Remove ignitio
F007 Recover or recycle if possible. Otherwise: incineration.
     Avoid electrostatic discharge generation.
```

```
Earth all equipment.
* *
     Avoid splash filling.
* *
     Use a vapour recovery system.
* *
     Keep in a well ventilated place.
* *
     Avoid naked flames. Remove ignition sources.
* *
     Do not smoke. Avoid sparks.
F008 HEDSET
F009 20-05-1994
F012 20
F020 119799
EOR
F002 27
F010 1.11
F004 3
F005 RM
F006 Transport
* *
* *
     UN number : 1120
* *
     Class : 3
* *
     Packing Group : III
* *
     Proper Shipping Name : Secondary butyl alcohol
* *
* *
     Sea (IMO)
* *
     Class: 3.3
* *
     Marine Pollutant : No
* *
     Symbol: Flammable liquid
* *
* *
     Rail/Road (ADR/RID)
* *
     Class : 3
* *
     Item : 31 c)
* *
     Symbol : Flammable
F007 Transport
* *
* *
     UN number : 1120
* *
     Class : 3
* *
     Packing Group : III
* *
     Proper Shipping Name : Secondary butyl alcohol
* *
     Sea (IMO)
* *
     Class : 3.3
* *
     Marine Pollutant : No
* *
     Symbol: Flammable liquid
* *
     Rail/Road (ADR/RID)
* *
     Class: 3
* *
     Item : 31 c)
* *
     Symbol : Flammable liquid
* *
     Kemler Plate : 30/1120
* *
* *
     Air (IATA/IACO)
* *
     Class : 3
* *
     Symbol: Flammable liquid
F008 HEDSET
F009 20-05-1994
F012 20
F020 119800
EOR
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F002 27
F010 1.11
F004 3
F005 SO
F006 SHELL FRANCE Rueil Malmaison
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 SHELL FRANCE Rueil Malmaison
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119801
EOR
F002 27
F010 1.11
F004 4
F005 RM
F006 Disposal options:
     Recover or recycle if possible. Otherwise: incineration.
* *
* *
     Transport classification:
* *
* *
     UN number: 1120
* *
     ADR/RID:
* *
     class/item: 3/31 c
* *
     packing group: 3
* *
     Kemler number: 30
* *
     label: 3
* *
     Proper shipping name: sec.-butyl alcohol
* *
* *
     ICAO:
* *
     С
F007 Disposal options:
* *
     Recover or recycle if possible. Otherwise: incineration.
* *
* *
     Transport classification:
* *
* *
     UN number: 1120
     ADR/RID:
* *
     class/item: 3/31 c
* *
     packing group: 3
* *
     Kemler number: 30
* *
     label: 3
* *
     Proper shipping name: sec.-butyl alcohol
* *
* *
     ICAO:
* *
     class: 3
* *
     packing group: III
* *
     label: flammable liquid
* *
     Proper shipping name: butanols
* *
* *
     IMO/IMDG:
* *
     class: 3.3
* *
     packing group: III
* *
     Marine pollutant: no
* *
     label: flammable liquid
* *
     IMDG page: 3313
```

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EMS number: 3-06
** MFAG plate: 305
F008 HEDSET
F009 30-05-1994
F012 20
F020 119802
EOR
F002 27
F010 1.11
F004 4
F005 S0
F006 Deutsche Shell Chemie GmbH Eschborn
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Deutsche Shell Chemie GmbH Eschborn
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119803
EOR
F002 27
F010 1.2
F004 1
F006 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119804
EOR
F002 27
F010 1.2
F004 2
F005 SO
F006 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119805
EOR
F002 27
F010 1.2
F004 3
F005 SO
F006 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
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F020 119806
EOR
F002 27
F010 1.2
F004 4
F005 SO
F006 Union Carbide Benelux Antwerpen
    EXXON CHEMICAL, Limited Fareham, Hampshire
* *
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
    Hamburg
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Union Carbide Benelux Antwerpen
** EXXON CHEMICAL, Limited Fareham, Hampshire
* *
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
    Hamburg
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119807
EOR
F002 27
F010 1.2
F004 5
F005 SO
F006 Union Carbide Benelux Antwerpen
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Union Carbide Benelux Antwerpen
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119808
EOR
F002 27
F010 1.2
F004 6
F005 SO
F006 Union Carbide Benelux Antwerpen
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Union Carbide Benelux Antwerpen
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119809
EOR
F002 27
F010 1.2
F004 7
F005 SO
F006 Atochem Paris la Defense
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Atochem Paris la Defense
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
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F012 20
F020 119810
EOR
F002 27
F010 1.2
F004 8
F005 SO
F006 SHELL FRANCE Rueil Malmaison
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 SHELL FRANCE Rueil Malmaison
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119811
EOR
F002 27
F010 1.2
F004 9
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
     Hamburg
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119812
EOR
F002 27
F010 1.2
F004 11
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
     Hamburg
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119813
EOR
F002 27
F010 1.2
F004 12
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
* *
     RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
     Hamburg
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EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
    Hamburg
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119814
EOR
F002 27
F010 1.2
F004 13
F005 SO
F006 Deutsche Shell Chemie GmbH Eschborn
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Deutsche Shell Chemie GmbH Eschborn
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119815
EOR
F002 27
F010 1.5
F004 8
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119816
EOR
F002 27
F010 1.6.1
F004 8
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119817
EOR
F002 27
F010 1.6.2
F004 13
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119818
EOR
F002 27
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F010 1.6.2
F004 14
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119819
EOR
F002 27
F010 1.6.2
F004 15
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119820
EOR
F002 27
F010 1.7
F004 37
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119821
EOR
F002 27
F010 1.7
F004 38
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119822
EOR
F002 27
F010 1.7
F004 39
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119823
EOR
F002 27
F010 1.7
F004 40
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F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119824
EOR
F002 27
F010 1.7
F004 41
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119825
EOR
F002 27
F010 1.7
F004 42
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119826
EOR
F002 27
F010 1.7
F004 43
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119827
EOR
F002 27
F010 1.7
F004 44
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119828
EOR
F002 27
F010 1.7
F004 45
F005 S0
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
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F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119829
EOR
F002 27
F010 1.7
F004 46
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119830
EOR
F002 27
F010 1.7
F004 47
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119831
EOR
F002 27
F010 1.7
F004 48
F005 SO
F006 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119832
EOR
F002 27
F010 1.8.1
F004 2
F005 RE
F006 Dutch MAC list 1994
F007 Dutch MAC list 1994
F008 HEDSET
F009 03-06-1994
F012 20
F020 119833
EOR
F002 27
F010 1.8.1
F004 2
F005 SO
F006 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
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EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119834
EOR
F002 27
F010 1.8.1
F004 3
F005 RE
F006 ACGIH list 1993-1994
F007 ACGIH list 1993-1994
F008 HEDSET
F009 03-06-1994
F012 20
F020 119835
EOR
F002 27
F010 1.8.1
F004 3
F005 SO
F006 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
** EUROPEAN COMMISSION - European Chemicals Bureau
                                                      Ispra (VA)
F007 Shell Nederland Chemie B.V. Hoogvliet-Rotterdam
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119836
EOR
F002 27
F010 1.8.1
F004 4
F005 RE
F006 1992-1993 Threshold Limit values and Biological Exposure
     Indices ACGIH
F007 1992-1993 Threshold Limit values and Biological Exposure
   Indices ACGIH
F008 HEDSET
F009 26-05-1994
F012 20
F020 119837
EOR
F002 27
F010 1.8.1
F004 4
F005 S0
F006 Union Carbide Benelux Antwerpen
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Union Carbide Benelux Antwerpen
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119838
EOR
F002 27
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F010 1.8.1
F004 5
F005 RE
F006 INRS, Valeurs limites d'exposition professionnelle aux
     substances dangereuses de l'ACGIH aux Etats-Unis et de la
* *
     Commission MAK en Allemagne, Cah. Notes Doc. 1992, 147,
* *
F007 INRS, Valeurs limites d'exposition professionnelle aux
     substances dangereuses de l'ACGIH aux Etats-Unis et de la
     Commission MAK en Allemagne, Cah. Notes Doc. 1992, 147,
     195-225
F008 HEDSET
F009 27-04-1994
F012 20
F020 119839
EOR
F002 27
F010 1.8.1
F004 5
F005 SO
F006 Atochem Paris la Defense
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Atochem Paris la Defense
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119840
EOR
F002 27
F010 1.8.1
F004 6
F005 RE
F006 INRS, Valeurs limites d'exposition professionnelle aux
     substances dangereuses de l'ACGIH aux Etats-Unis et de la
     Commission MAK en Allemagne, Cah. Notes Doc. 1992, 147,
* *
     195-225
F007 INRS, Valeurs limites d'exposition professionnelle aux
     substances dangereuses de l'ACGIH aux Etats-Unis et de la
* *
     Commission MAK en Allemagne, Cah. Notes Doc. 1992, 147,
* *
     195-225
F008 HEDSET
F009 27-04-1994
F012 20
F020 119841
EOR
F002 27
F010 1.8.1
F004 6
F005 SO
F006 Atochem Paris la Defense
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Atochem Paris la Defense
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
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F020 119842
EOR
F002 27
F010 1.8.1
F004 7
F005 CT
F006 France
F007 France
F008 HEDSET
F009 27-04-1994
F012 20
F020 119843
EOR
F002 27
F010 1.8.1
F004 7
F005 RE
F006 INRS, Valeurs limites d'exposition professionnelle aux
     agents chimiques en France, Cah. Notes Doc. 1993, 153,
* *
F007 INRS, Valeurs limites d'exposition professionnelle aux
     agents chimiques en France, Cah. Notes Doc. 1993, 153,
* *
     557-574
F008 HEDSET
F009 27-04-1994
F012 20
F020 119844
EOR
F002 27
F010 1.8.1
F004 7
F005 SO
F006 Atochem Paris la Defense
   EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Atochem Paris la Defense
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119845
EOR
F002 27
F010 1.8.1
F004 8
F005 RE
F006 Threshold Limit Values Book 1993-1994.
F007 Threshold Limit Values Book 1993-1994.
F008 HEDSET
F009 26-05-1994
F012 20
F020 119846
EOR
F002 27
F010 1.8.1
F004 8
F005 RM
F006 Use local exhaust ventilation.
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* *
     Hand protection: PVC, nitrile or neoprene gloves.
* *
     Eye protection : safety monogoggles.
* *
     Body protection: standard issue work clothes.
* *
                       chemicals resistant safety shoes or boots.
F007 Use local exhaust ventilation.
* *
     Hand protection: PVC, nitrile or neoprene gloves.
     Eye protection : safety monogoggles.
* *
* *
     Body protection : standard issue work clothes.
                       chemicals resistant safety shoes or boots.
F008 HEDSET
F009 26-05-1994
F012 20
F020 119847
EOR
F002 27
F010 1.8.1
F004 8
F005 SO
F006 SHELL FRANCE Rueil Malmaison
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 SHELL FRANCE Rueil Malmaison
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119848
EOR
F002 27
F010 1.8.1
F004 9
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
* *
     RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
     Hamburg
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
     Hamburg
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119849
EOR
F002 27
F010 1.8.1
F004 10
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
     Hamburg
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
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F009 09-01-2002
F012 20
F020 119850
EOR
F002 27
F010 1.8.1
F004 11
F005 S0
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119851
EOR
F002 27
F010 1.8.1
F004 12
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
    Hamburg
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
* *
    Hamburg
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119852
EOR
F002 27
F010 1.8.1
F004 13
F005 RM
F006 Substance is easily resorbed.
F007 Substance is easily resorbed.
F008 HEDSET
F009 30-05-1994
F012 20
F020 119853
EOR
F002 27
F010 1.8.1
F004 13
F005 SO
F006 Deutsche Shell Chemie GmbH Eschborn
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 Deutsche Shell Chemie GmbH Eschborn
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119854
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EOR
F002 27
F010 1.8.1
F004 16
F005 SO
F006 RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 RWE-DEA Aktiengesellschaft für Mineraloel und Chemie
    Hamburg
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119855
EOR
F002 27
F010 2.1
F004 1
F005 RE
F006 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
     Reinhold Company, New York, NY, USA.
F007 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
     Reinhold Company, New York, NY, USA.
F008 CLGETTS
F012 20
F020 119862
EOR
F002 27
F010 2.1
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119863
EOR
F002 27
F010 2.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119864
EOR
F002 27
F010 2.1
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F004 2
F005 RE
F006 Hawley's Condensed Chemical Dictionary. Lewis, R. J., Sr., ed. 13th ed.
     New York, NY: John Wiley & Sons, Inc. 1997.
F007 Hawley's Condensed Chemical Dictionary. Lewis, R. J., Sr., ed. 13th ed.
     New York, NY: John Wiley & Sons, Inc. 1997.
F008 CLGETTS
F012 20
F020 119865
EOR
F002 27
F010 2.1
F004 2
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119866
EOR
F002 27
F010 2.1
F004 2
F005 S0
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119867
EOR
F002 27
F010 2.10
F004 1
F005 RE
F006 Safety Data Sheet (sec-butanol), June 1990, ELF ATOCHEM.
F007 Safety Data Sheet (sec-butanol), June 1990, ELF ATOCHEM.
F008 HEDSET
F009 02-06-1994
F012 20
F020 119868
EOR
F002 27
F010 2.10
F004 1
F005 RM
F006 Explosive limits of vapours in air: 1.7 to 9.8% vol.
     Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Explosive limits of vapours in air: 1.7 to 9.8% vol.
     Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F009 02-06-1994
F012 20
F020 119869
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EOR
F002 27
F010 2.10
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119870
EOR
F002 27
F010 2.2
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119871
EOR
F002 27
F010 2.2
F004 1
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119872
EOR
F002 27
F010 2.2
F004 2
F005 RE
F006 Ethel Browning's Toxicity and Metabolism of Industrial Solvents. Snyder,
     R., ed. Second Edition. Volume 3 Alcohols and Esters. New York, NY:
     Elsevier, 1992.
F007 Ethel Browning's Toxicity and Metabolism of Industrial Solvents. Snyder,
     R., ed. Second Edition. Volume 3 Alcohols and Esters. New York, NY:
     Elsevier, 1992.
F008 CLGETTS
F012 20
F020 119873
EOR
F002 27
F010 2.2
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F004 2
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119874
EOR
F002 27
F010 2.2
F004 2
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119875
EOR
F002 27
F010 2.3
F004 1
F005 RE
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
F008 CLGETTS
F012 20
F020 119876
EOR
F002 27
F010 2.3
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
   EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119877
EOR
F002 27
F010 2.3
F004 2
F005 RE
F006 Safety Data Sheet ELF ATOCHEM June 1990;
F007 Safety Data Sheet ELF ATOCHEM June 1990;
F008 HEDSET
F009 02-06-1994
F012 20
F020 119878
EOR
F002 27
F010 2.3
F004 2
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F005 RM
F006 Vapour density
F007 Vapour density
F008 HEDSET
F009 02-06-1994
F012 20
F020 119879
EOR
F002 27
F010 2.3
F004 2
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119880
EOR
F002 27
F010 2.3
F004 3
F005 RE
F006 CRC Handbook of Chemistry and Physics. Lide, D. R., ed. 81st Edition,
    pp. 3-100. CRC Press LLC, Boca Raton, FL. 2000.
F007 CRC Handbook of Chemistry and Physics. Lide, D. R., ed. 81st Edition,
     pp. 3-100. CRC Press LLC, Boca Raton, FL. 2000.
F008 CLGETTS
F012 20
F020 119881
EOR
F002 27
F010 2.3
F004 3
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119882
EOR
F002 27
F010 2.3
F004 3
F005 SO
F006 ExxonBiomedical Sciences, Inc., New Jersey, USA
F007 ExxonBiomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119883
EOR
F002 27
F010 2.4
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F004 1
F005 RE
F006 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
     Reinhold Company, New York, NY, USA.
F007 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
     Reinhold Company, New York, NY, USA.
F008 CLGETTS
F012 20
F020 119884
EOR
F002 27
F010 2.4
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119885
EOR
F002 27
F010 2.4
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119886
EOR
F002 27
F010 2.5
F004 2
F005 RE
F006 Hansch, C., Leo, A., Hoekman, D. 1995. Exploring QSAR - Hydrophobic,
     Electronic, and Steric Constants. American Chemical Society, Washington,
     DC, USA.
F007 Hansch, C., Leo, A., Hoekman, D. 1995. Exploring QSAR - Hydrophobic,
     Electronic, and Steric Constants. American Chemical Society, Washington,
     DC, USA.
F008 CLGETTS
F012 20
F020 119887
EOR
F002 27
F010 2.5
F004 2
F005 RM
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F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119888
EOR
F002 27
F010 2.5
F004 2
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119889
EOR
F002 27
F010 2.6.1
F004 1
F005 RE
F006 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
    Reinhold Company, New York, NY, USA.
F007 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
     Reinhold Company, New York, NY, USA.
F008 CLGETTS
F012 20
F020 119890
EOR
F002 27
F010 2.6.1
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119891
EOR
F002 27
F010 2.6.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
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    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
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F020 119892
EOR
F002 27
F010 2.6.1
F004 2
F005 RE
F006 Handbook of Environmental Data on Organic Chemicals, Second
     Edition. Karel Verschueren, ed., Van Nostrand Reinhold
     Company (1983), pp. 301-302.
F007 Handbook of Environmental Data on Organic Chemicals, Second
    Edition. Karel Verschueren, ed., Van Nostrand Reinhold
     Company (1983), pp. 301-302.
F008 HEDSET
F009 02-06-1994
F012 20
F020 119893
EOR
F002 27
F010 2.6.1
F004 2
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119894
EOR
F002 27
F010 2.6.1
F004 2
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119895
EOR
F002 27
F010 2.6.1
F004 3
F006 Hefter, G.T. 1984. Solub. Data Ser., 15:94-119.
F007 Hefter, G.T. 1984. Solub. Data Ser., 15:94-119.
F008 CLGETTS
F012 20
F020 119896
EOR
F002 27
F010 2.6.1
F004 3
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F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119897
EOR
F002 27
F010 2.6.1
F004 3
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119898
EOR
F002 27
F010 2.7
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
   EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119899
EOR
F002 27
F010 2.7
F004 2
F005 RE
F006 Safety Data Sheet (sec-butanol), June 1990, ELF ATOCHEM.
F007 Safety Data Sheet (sec-butanol), June 1990, ELF ATOCHEM.
F008 HEDSET
F009 02-06-1994
F012 20
F020 119900
EOR
F002 27
F010 2.7
F004 2
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119901
EOR
F002 27
F010 2.7
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F004 2
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119902
EOR
F002 27
F010 2.8
F004 1
F005 RE
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
F008 CLGETTS
F012 20
F020 119903
EOR
F002 27
F010 2.8
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119904
EOR
F002 27
F010 2.8
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119905
EOR
F002 27
F010 2.8
F004 2
F005 RE
F006 Safety Data Sheet ELF ATOCHEM June 1990
F007 Safety Data Sheet ELF ATOCHEM June 1990
F008 HEDSET
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F009 02-06-1994
F012 20
F020 119906
EOR
F002 27
F010 2.8
F004 2
F005 S0
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119907
EOR
F002 27
F010 2.9
F004 1
F005 RE
F006 Safety Data Sheet (sec-butanol), June 1990, ELF ATOCHEM.
F007 Safety Data Sheet (sec-butanol), June 1990, ELF ATOCHEM.
F008 HEDSET
F009 02-06-1994
F012 20
F020 119908
F:OR
F002 27
F010 2.9
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
    commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119909
EOR
F002 27
F010 2.9
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
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     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119910
EOR
F002 27
F010 3.1.1
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F004 1
F005 RE
F006 Chemcial Hazard Assessment Division, Syracuse Research Corp. 1988.
     Syracuse, NY, USA.
F007 Chemcial Hazard Assessment Division, Syracuse Research Corp. 1988.
     Syracuse, NY, USA.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119911
EOR
F002 27
F010 3.1.1
F004 1
F005 RM
F006 2-Butanol does not contain chromophores that adsorb light at wavelengths
     >290 nm. Therefore, direct photolysis will not occur.
F007 2-Butanol does not contain chromophores that adsorb light at wavelengths
     >290 nm. Therefore, direct photolysis will not occur.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119912
EOR
F002 27
F010 3.1.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119913
EOR
F002 27
F010 3.1.1
F004 4
F005 RE
F006 Dilling, W. L. et. al. (1976). Organic Photochemistry.
     Simulated atmospheric photodecomposition rates of methylene
* *
     chloride, 1,1,1-trichloroethane, trichloroethylene,
* *
     tetrachloroethylene, and other compounds. Env. Sci.
* *
     Technol. 10(4)
F007 Dilling, W. L. et. al. (1976). Organic Photochemistry.
* *
     Simulated atmospheric photodecomposition rates of methylene
* *
     chloride, 1,1,1-trichloroethane, trichloroethylene,
* *
     tetrachloroethylene, and other compounds. Env. Sci.
* *
     Technol. 10(4), 351-356.
F008 HEDSET
F009 09-05-1994
F012 20
F020 119914
EOR
F002 27
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F010 3.1.1
F004 4
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119915
EOR
F002 27
F010 3.1.1
F004 4
F005 RM
F006 Under simulated atmospheric conditions and in the presence
     of 5 ppm NO, the half life of 2-butanol (10 ppm) is 4 hours.
F007 Under simulated atmospheric conditions and in the presence
     of 5 ppm NO, the half life of 2-butanol (10 ppm) is 4 hours.
F008 HEDSET
F009 09-05-1994
F012 20
F020 119916
EOR
F002 27
F010 3.1.1
F004 4
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119917
EOR
F002 27
F010 3.1.1
F004 5
F005 RE
F006 Shell International Petroleum, B.V.
F007 Shell International Petroleum, B.V.
F008 CLGETTS
F012 20
F020 119918
EOR
F002 27
F010 3.1.1
F004 5
F005 RM
F006 Calculated value using draft OECD method of Atkinson (1990).
F007 Calculated value using draft OECD method of Atkinson (1990).
F008 CLGETTS
F012 20
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F020 119919
EOR
F002 27
F010 3.1.1
F004 5
F005 S0
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119920
EOR
F002 27
F010 3.1.1
F004 6
F005 RE
F006 EPIWIN. 1999. Estimation Program Interface for Windows, Version 3.04.
     Syracuse Research Corporation, Syracuse, NY, USA.
F007 EPIWIN. 1999. Estimation Program Interface for Windows, Version 3.04.
     Syracuse Research Corporation, Syracuse, NY, USA.
F008 CLGETTS
F012 20
F020 119921
EOR
F002 27
F010 3.1.1
F004 6
F005 RL
F006 Rated a 2 for reliability because it is a calculated value.
F007 Rated a 2 for reliability because it is a calculated value.
F008 CLGETTS
F012 20
F020 119922
EOR
F002 27
F010 3.1.1
F004 6
F005 RM
F006 50% degradation after 1.07 days based on a 12-hr. day. The SMILES
     (Simplified Molecular Input Line Entry System) structure used with the
     model was: OC(CC)C.
F007 50% degradation after 1.07 days based on a 12-hr. day. The SMILES
     (Simplified Molecular Input Line Entry System) structure used with the
     model was: OC(CC)C.
F008 CLGETTS
F012 20
F020 119923
EOR
F002 27
F010 3.1.1
F004 6
F005 RM
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F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119924
EOR
F002 27
F010 3.1.1
F004 6
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119925
EOR
F002 27
F010 3.1.1
F004 7
F005 RE
F006 Edney, E.O. and Corse, E.W. 1986. Hydroxyl radical rate constant
     intercomparison study. EPA/600/3-86/056, US EPA, Research Triangle Park,
    NC, US.
F007 Edney, E.O. and Corse, E.W. 1986. Hydroxyl radical rate constant
     intercomparison study. EPA/600/3-86/056, US EPA, Research Triangle Park,
    NC, US.
F008 CLGETTS
F012 20
F020 119926
EOR
F002 27
F010 3.1.1
F004 7
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119927
EOR
F002 27
F010 3.1.1
F004 7
F005 RM
F006 Rate constants were determined from data developed in three labs. Two
     equations were applied to the measured data and designated the TI
     (time-included) and TE (time-excluded) methods. Both equations are in the
     form of a straight line equati
F007 Rate constants were determined from data developed in three labs. Two
     equations were applied to the measured data and designated the TI
     (time-included) and TE (time-excluded) methods. Both equations are in the
     form of a straight line equation, y=mx+b. Least square linear regression
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analyses were applied to the data from each experiment.

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F008 CLGETTS
F012 20
F020 119928
EOR
F002 27
F010 3.1.1
F004 7
F005 RS
F006 Hydroxyl rate constants as calculated from measured data using the
     time-included equation:
     10.30 +/- 2.48 \times 10-12 \text{ cm}3/\text{molecule-sec (r2 = 0.97); @ 24C}
* *
     11.55 + - 1.77 \times 10-12 \text{ cm}^3/\text{molecule-sec (r2 = 0.99); @ 24C}
* *
     4.01 +/- 1.38 \times 10-12 \text{ cm3/mole}
F007 Hydroxyl rate constants as calculated from measured data using the
     time-included equation:
* *
* *
     10.30 + - 2.48 \times 10-12 \text{ cm}3/\text{molecule-sec} (r2 = 0.97); @ 24C
* *
     11.55 + - 1.77 \times 10-12 \text{ cm}^3/\text{molecule-sec (r2 = 0.99); @ 24C}
* *
     4.01 + - 1.38 \times 10-12 \text{ cm}3/\text{molecule-sec} (r2 = 0.76); @ 34C
* *
     Hydroxyl rate constants as calculated from measured data using the
     time-excluded equation:
* *
     9.40 + - 0.87 \times 10-12 \text{ cm}3/\text{molecule-sec} (r2 = 0.99); @ 24C
* *
     7.37 + -1.65 \times 10-12 \text{ cm}^3/\text{molecule-sec (r2} = 0.99); @ 24C
* *
     2.71 + - 0.20 \times 10 - 12 \text{ cm}^3/\text{molecule-sec (r2} = 0.97); @ 34C
F008 CLGETTS
F012 20
F020 119929
EOR
F002 27
F010 3.1.1
F004 7
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EXXON CHEMICAL, Limited Fareham, Hampshire
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F012 20
F020 119930
EOR
F002 27
F010 3.1.2
F004 2
F005 RE
F006 Harris, J.C. 1982. Rate of Hydrolysis. In: Handbook of Chemical Property
     Estimation Methods. Environmental Behavior of Organic Compounds. 7, 1-48.
     Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds). McGraw-Hill, New
     York, NY, USA.
F007 Harris, J.C. 1982. Rate of Hydrolysis. In: Handbook of Chemical Property
     Estimation Methods. Environmental Behavior of Organic Compounds. 7, 1-48.
     Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds). McGraw-Hill, New
     York, NY, USA.
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F008 HEDSET
F009 15-02-1994
F012 20
F020 119931
EOR
F002 27
F010 3.1.2
F004 2
F005 RS
F006 Hydrolysis will not contribute to the transformation of sBA in aquatic
     environments. Hydrolysis of an organic chemical is the transformation
     process in which a water molecule or hydroxide ion reacts to form a new
     carbon-oxygen bond, thereb
F007 Hydrolysis will not contribute to the transformation of sBA in aquatic
     environments. Hydrolysis of an organic chemical is the transformation
    process in which a water molecule or hydroxide ion reacts to form a new
     carbon-oxygen bond, thereby changing the parent chemical. Chemicals that
    are susceptible to hydrolysis contain functional groups that can be
    displaced by a nucleophilic substitution reaction. Simple alcohols such
     as sBA are resistant to hydrolysis because they lack a functional group
     that is hydrolytically reactive.
F008 HEDSET
F009 15-02-1994
F012 20
F020 119932
EOR
F002 27
F010 3.1.2
F004 2
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119933
EOR
F002 27
F010 3.1.3
F004 1
F005 RE
F006 Lyman, W.J. (1982). Adsorption Coefficient for Soils and Sediments. In:
     Handbook of Chemical Property Estimation Methods. Environmental Behavior
     of Organic Compounds. 4,1-33. Lyman W.J., Reehl, W.F., and Rosenblatt,
     D.H. (eds). McGraw-Hill,
F007 Lyman, W.J. (1982). Adsorption Coefficient for Soils and Sediments. In:
     Handbook of Chemical Property Estimation Methods. Environmental Behavior
     of Organic Compounds. 4,1-33. Lyman W.J., Reehl, W.F., and Rosenblatt,
     D.H. (eds). McGraw-Hill, New York, NY, USA.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119934
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EOR
F002 27
F010 3.1.3
F004 1
F005 RE
F006 Thomas, R.G. 1982. Volatilzation from Soil. In: Handbook of Chemical
     Property Estimation Methods. Environmental Behavior of Organic Compounds.
     16, 1-50. Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds).
     McGraw-Hill, New York, NY, USA.
F007 Thomas, R.G. 1982. Volatilzation from Soil. In: Handbook of Chemical
     Property Estimation Methods. Environmental Behavior of Organic Compounds.
     16, 1-50. Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds).
     McGraw-Hill, New York, NY, USA.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119935
EOR
F002 27
F010 3.1.3
F004 1
F005 RM
F006 If released on land, 2-butanol has the potential to leach
     into soil. 2-Butanol also has the potential to volatilize from dry soil.
F007 If released on land, 2-butanol has the potential to leach
     into soil. 2-Butanol also has the potential to volatilize from dry soil.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119936
EOR
F002 27
F010 3.1.3
F004 1
F005 RM
F006 The estimated log Koc for 2-butanol is 0.84. Therefore,
* *
     2-butanol should not absorb significantly to soil or
* *
     sediment.
     The Koc value was calculated using the equation in Lyman (1982): log Koc
     = -0.55\log S + 3.64 (S, water solubility, in
F007 The estimated log Koc for 2-butanol is 0.84. Therefore,
     2-butanol should not absorb significantly to soil or
* *
     sediment.
     The Koc value was calculated using the equation in Lyman (1982): log Koc
     = -0.55\log S + 3.64 (S, water solubility, in mg/L); water solubility
     value used = 125,000 \text{ mg/L} from Hahn et al. (1986)
* *
     [Lyman, W.J. (1982). Adsorption Coefficient for Soils and Sediments. In:
     Handbook of Chemical Property Estimation Methods. Environmental Behavior
     of Organic Compounds. 4,1-33. Lyman W.J., Reehl, W.F., and Rosenblatt,
     D.H. (eds). McGraw-Hill, New York, NY, USA.]
* *
     [Hahn, H-D., Dämbkes, G., and Rupprich, N. (1986). Butanols. In: Gerhartz
     W (ed), Ullmann's encyclopedia of industrial chemistry, 5th ed, vol A4,
     benzyl alcohol to calcium sulfate. VCH, Weinheim, 463-474.]
F008 HEDSET
F009 21-10-1992
F012 20
F020 119937
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EOR
F002 27
F010 3.1.3
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119938
EOR
F002 27
F010 3.2.1
F004 1
F005 RE
F006 Lucas, S.V. 1984. GC/MS Analysis of organics in drinking water
     concentrates and advanced treatment concentrates. Vol. 1.
     USEPA-600/1-84-020a (NTIS).
F007 Lucas, S.V. 1984. GC/MS Analysis of organics in drinking water
     concentrates and advanced treatment concentrates. Vol. 1.
     USEPA-600/1-84-020a (NTIS).
F008 HEDSET
F009 21-10-1992
F012 20
F020 119939
EOR
F002 27
F010 3.2.1
F004 1
F005 RM
F006 2-Butanol was identified in drinking water from Ottumwa, IA.
F007 2-Butanol was identified in drinking water from Ottumwa, IA.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119940
EOR
F002 27
F010 3.2.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119941
EOR
F002 27
F010 3.2.1
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F004 2
F005 RE
F006 Great Lakes Water Quality Board. 1983. Inventory of Chemical
     Substances Indentified in the Great Lakes Ecosystem. Report
     to the Great Lakes Water Quality Board, Windsor, Canada.
F007 Great Lakes Water Quality Board. 1983. Inventory of Chemical
     Substances Indentified in the Great Lakes Ecosystem. Report
     to the Great Lakes Water Quality Board, Windsor, Canada.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119942
EOR
F002 27
F010 3.2.1
F004 2
F005 RM
F006 2-Butanol has been detected in the Niagara River (Lake
     Ontario basin), but not in the western basin of Lake
F007 2-Butanol has been detected in the Niagara River (Lake
     Ontario basin), but not in the western basin of Lake
* *
     Ontario.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119943
F:OR
F002 27
F010 3.2.1
F004 2
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119944
EOR
F002 27
F010 3.2.1
F004 3
F005 RE
F006 Shackelford, W.M. et al. 1983. Analyt. Chim. Acta, 146:15-27
     (supplemental data).
F007 Shackelford, W.M. et al. 1983. Analyt. Chim. Acta, 146:15-27
     (supplemental data).
F008 HEDSET
F009 21-10-1992
F012 20
F020 119945
EOR
F002 27
F010 3.2.1
F004 3
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F005 RM
F006 In a comprehensive survey of wastewater from 4000 industrial and
     publically owned treatment works sponsored by the Effluent Guidelines
     Division of the U.S. EPA, 2-butanol was identified in discharges of the
     following industrial category (po
F007 In a comprehensive survey of wastewater from 4000 industrial and
    publically owned treatment works sponsored by the Effluent Guidelines
    Division of the U.S. EPA, 2-butanol was identified in discharges of the
     following industrial category (positive occurences, median concentration
     in ppb): leather tanning (1; 46.7), petroleum refining (1; 149.3), paint
     and ink (1; 324.7), organics and plastics (2; 35.4), pesticides
    manufacture (1; 36.2), pharmaceuticals (1; 4.6), foundries (1; 41.0),
     electronics (1; 19.9), mechanical products (3; 90.6), publically owned
     treatment works (3; 12.4). The highest effluent concentration was 920.1
     ppb in the mechanical products industry.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119946
EOR
F002 27
F010 3.2.1
F004 3
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119947
EOR
F002 27
F010 3.2.1
F004 4
F005 RE
F006 Sawhney, B.L. and Kozloski, R.P. 1984. J. Environ. Qual. 13:349-352.
F007 Sawhney, B.L. and Kozloski, R.P. 1984. J. Environ. Qual. 13:349-352.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119948
EOR
F002 27
F010 3.2.1
F004 4
F006 2-Butanol was found in landfill leachate from 1 of 5 sites
     in Connecticut. The concentration in this leachate ranged
     from 6.2 to 14.9 ppm.
F007 2-Butanol was found in landfill leachate from 1 of 5 sites
    in Connecticut. The concentration in this leachate ranged
* *
    from 6.2 to 14.9 ppm.
F008 HEDSET
F009 21-10-1992
F012 20
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F020 119949
EOR
F002 27
F010 3.2.1
F004 4
F005 S0
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119950
EOR
F002 27
F010 3.2.1
F004 5
F005 RE
F006 Juttner, F. 1986. Chemosphere, 15: 985-992.
F007 Juttner, F. 1986. Chemosphere, 15: 985-992.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119951
EOR
F002 27
F010 3.2.1
F004 5
F005 RM
F006 2-Butanol was detected but not quantified in forest air in
     the southern Black Forest of Germany.
F007 2-Butanol was detected but not quantified in forest air in
    the southern Black Forest of Germany.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119952
EOR
F002 27
F010 3.2.1
F004 5
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119953
EOR
F002 27
F010 3.2.1
F004 6
F005 RE
F006 Snider, J.R. and Dawson, G.A. 1985. J. Geophys. Res., 90:3797-3805.
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F007 Snider, J.R. and Dawson, G.A. 1985. J. Geophys. Res., 90:3797-3805.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119954
EOR
F002 27
F010 3.2.1
F004 6
F005 RM
F006 Low molecular weight alcohols, including 2-butanol, were not detected in
     air or precipitation samples taken at Tucson, Arizona, and two rural
     sites 40 km away.
F007 Low molecular weight alcohols, including 2-butanol, were not detected in
     air or precipitation samples taken at Tucson, Arizona, and two rural
     sites 40 km away.
F008 HEDSET
F009 21-10-1992
F012 20
F020 119955
EOR
F002 27
F010 3.2.1
F004 6
F005 S0
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119956
EOR
F002 27
F010 3.3.1
F004 1
F005 RE
F006 Thomas, R.G. 1982. Volatilization from Water. In: Handbook of Chemical
     Property Estimation Methods. Environmental Behavior of Organic Compounds.
     15, 1-34. Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds).
     McGraw-Hill, New York, NY, USA.
F007 Thomas, R.G. 1982. Volatilization from Water. In: Handbook of Chemical
     Property Estimation Methods. Environmental Behavior of Organic Compounds.
     15, 1-34. Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds).
     McGraw-Hill, New York, NY, USA.
F008 HEDSET
F009 10-05-1994
F012 20
F020 119957
EOR
F002 27
F010 3.3.1
F004 1
F005 RS
F006 On the basis of Henry's Law constant, the volatilization half-life of
     2-butanol is estimated to be 120.2 hours in a model river with the
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follwoing parameters: 1 m deep, 1 m/sec flow rate, and 3 m/sec wind speed.
F007 On the basis of Henry's Law constant, the volatilization half-life of
     2-butanol is estimated to be 120.2 hours in a model river with the
     follwoing parameters: 1 m deep, 1 m/sec flow rate, and 3 m/sec wind speed.
F008 HEDSET
F009 10-05-1994
F012 20
F020 119958
EOR
F002 27
F010 3.3.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119959
EOR
F002 27
F010 3.3.2
F004 1
F005 RE
F006 Mackay, D. 1998. Level I Fugacity-Based Environmental Equilibrium
     Partitioning Model, Version 2.1 (16-bit). Environmental Modelling Centre,
     Trent University, Ontario, Canada.
F007 Mackay, D. 1998. Level I Fugacity-Based Environmental Equilibrium
     Partitioning Model, Version 2.1 (16-bit). Environmental Modelling Centre,
     Trent University, Ontario, Canada.
F008 HEDSET
F009 10-05-1994
F012 20
F020 119960
EOR
F002 27
F010 3.3.2
F004 1
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 HEDSET
F012 20
F020 119961
EOR
F002 27
F010 3.3.2
F004 1
F005 RM
F006 Physicochemical data used in the calculation:
* *
* *
                           Value w/ Units
     Parameter
* *
```

```
Molecular Weight
                            74.12
* *
                            20 C
     Temperature
* *
     Log Kow
                            0.61
* *
     Water Solubility
                            125,000 g/m3
* *
     Vapor Pressure
                            1600 Pa
* *
     Melting
F007 Physicochemical data used in the calculation:
* *
     Parameter
                            Value w/ Units
* *
* *
    Molecular Weight
                            74.12
* *
    Temperature
                            20 C
* *
   Log Kow
                            0.61
* *
    Water Solubility
                            125,000 g/m3
* *
     Vapor Pressure
                            1600 Pa
* *
    Melting Point
                            -114 C
F008 HEDSET
F009 10-05-1994
F012 20
F020 119962
EOR
F002 27
F010 3.3.2
F004 1
F005 RS
F006 Using the Mackay Level I calculation, the following
     distribution is predicted for 2-butanol:
* *
* *
     %Distribution
                      Compartment
* *
* *
       16.28
                          Air
* *
       83.68
                          Water
* *
       0.03
                          Soil
* *
        0.01
                          Sediment
* *
        0.00
F007 Using the Mackay Level I calculation, the following
* *
     distribution is predicted for 2-butanol:
* *
* *
    %Distribution
                      Compartment
* *
* *
       16.28
                          Air
* *
       83.68
                          Water
* *
       0.03
                          Soil
* *
        0.01
                          Sediment
* *
        0.00
                         Suspended Sediment
* *
        0.00
                         Biota
F008 HEDSET
F012 20
F020 119963
EOR
F002 27
F010 3.3.2
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
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F007 EXXON CHEMICAL, Limited Fareham, Hampshire
** ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119964
EOR
F002 27
F010 3.5
F004 2
F005 RE
F006 Sasaki, S. 1978. pp 283-298 in Aquatic Pollutants: Transfer
     and biological effects. Hutzinger, O. et. al. (eds.)
     Pergamon Press.
F007 Sasaki, S. 1978. pp 283-298 in Aquatic Pollutants: Transfer
     and biological effects. Hutzinger, O. et. al. (eds.)
     Pergamon Press.
F008 HEDSET
F009 15-02-1994
F012 20
F020 119965
EOR
F002 27
F010 3.5
F004 2
F005 RM
F006 98% BOD reduction after 5 days
F007 98% BOD reduction after 5 days
F008 HEDSET
F009 15-02-1994
F012 20
F020 119966
EOR
F002 27
F010 3.5
F004 2
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119967
EOR
F002 27
F010 3.5
F004 3
F005 RE
F006 Yonezawa, Y. and Urushigawa, Y. 1979. Chemosphere 8,
    139-142.
F007 Yonezawa, Y. and Urushigawa, Y. 1979. Chemosphere 8,
     139-142.
F008 HEDSET
F009 15-02-1994
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F012 20
F020 119968
EOR
F002 27
F010 3.5
F004 3
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119969
EOR
F002 27
F010 3.5
F004 4
F005 RE
F006 Chou, W.L. et.al. 1979. Bio. Eng. Symp. 8, 391-414.
F007 Chou, W.L. et.al. 1979. Bio. Eng. Symp. 8, 391-414.
F008 HEDSET
F009 15-02-1994
F012 20
F020 119970
EOR
F002 27
F010 3.5
F004 4
F005 RM
F006 In a long-term study using anaerobic upflow filters, 93%
    utilization rate was seen after a 52 day operation.
F007 In a long-term study using anaerobic upflow filters, 93%
    utilization rate was seen after a 52 day operation.
F008 HEDSET
F009 15-02-1994
F012 20
F020 119971
EOR
F002 27
F010 3.5
F004 4
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119972
EOR
F002 27
F010 3.5
F004 5
F005 RE
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F006 Dore et al. 1975. Trib. Cebedeau, 27:3-11.
F007 Dore et al. 1975. Trib. Cebedeau, 27:3-11.
F008 HEDSET
F009 10-05-1994
F012 20
F020 119973
EOR
F002 27
F010 3.5
F004 5
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119974
EOR
F002 27
F010 3.5
F004 5
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119975
EOR
F002 27
F010 3.5
F004 7
F005 RE
F006 Wagner. 1974. Wasser, 42:271-305.
F007 Wagner. 1974. Wasser, 42:271-305.
F008 HEDSET
F009 10-05-1994
F012 20
F020 119976
EOR
F002 27
F010 3.5
F004 7
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119977
EOR
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F002 27
F010 3.5
F004 7
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119978
EOR
F002 27
F010 3.5
F004 8
F005 RE
F006 Gerhold and Malaney. 1966. J. Water Pollution Control Fed., 38:562-579.
F007 Gerhold and Malaney. 1966. J. Water Pollution Control Fed., 38:562-579.
F008 HEDSET
F009 10-05-1994
F012 20
F020 119979
EOR
F002 27
F010 3.5
F004 8
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119980
EOR
F002 27
F010 3.5
F004 8
F005 S0
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119981
EOR
F002 27
F010 3.5
F004 9
F005 RE
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F006 Pitter. 1976. Water Research, 10:231-235.
F007 Pitter. 1976. Water Research, 10:231-235.
F008 HEDSET
F009 10-05-1994
F012 20
F020 119982
EOR
F002 27
F010 3.5
F004 9
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 119983
EOR
F002 27
F010 3.5
F004 9
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 119984
EOR
F002 27
F010 3.5
F004 10
F005 ME
F006 The APHA No. 219 test method, which measures dissolved oxygen, is similar
     to the OECD 301D, Closed Bottle biodegradation test procedure.
F007 The APHA No. 219 test method, which measures dissolved oxygen, is similar
     to the OECD 301D, Closed Bottle biodegradation test procedure.
F008 CLGETTS
F012 20
F020 119985
EOR
F002 27
F010 3.5
F004 10
F005 RE
F006 Bridie, A.L., C.J.M. Wolff, and M. Winter. 1979. BOD and COD of Some
     Petrochemicals. Water Research. 13:627-630.
F007 Bridie, A.L., C.J.M. Wolff, and M. Winter. 1979. BOD and COD of Some
     Petrochemicals. Water Research. 13:627-630.
F008 CLGETTS
F012 20
F020 119986
```

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EOR
F002 27
F010 3.5
F004 10
F005 RL
F006 Although it was cited that a standard method was followed, there is
     little information in the article confirming that the test was conducted
     according to the method description. This lack of information supports a
    reliability rating of 2.
F007 Although it was cited that a standard method was followed, there is
     little information in the article confirming that the test was conducted
     according to the method description. This lack of information supports a
     reliability rating of 2.
F008 CLGETTS
F012 20
F020 119987
EOR
F002 27
F010 3.5
F004 10
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
    commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119988
EOR
F002 27
F010 3.5
F004 10
F005 RS
F006 The ThOD (theoretical oxygen demand) = 2.59 g/g. The measured BOD
     (biological oxygen demand) = 2.15 g/g.
     The only deviation to the test method was that 0.5 mg/L of allylthiourea
     was added to the test medium to prevent nitrification.
F007 The ThOD (theoretical oxygen demand) = 2.59 g/g. The measured BOD
     (biological oxygen demand) = 2.15 g/g.
    The only deviation to the test method was that 0.5 mg/L of allylthiourea
     was added to the test medium to prevent nitrification.
F008 CLGETTS
F012 20
F020 119989
EOR
F002 27
F010 3.5
F004 10
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119990
EOR
F002 27
F010 3.5
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F004 10
F005 TC
F006 10 ml of filtered wastewater was used as an inoculum with 500 ml of test
     medium. The source of the inoculum from within the wastewater treatment
     plant was not stated. The inoculum was not acclimated. There was no
     information on controls and
F007 10 ml of filtered wastewater was used as an inoculum with 500 ml of test
    medium. The source of the inoculum from within the wastewater treatment
    plant was not stated. The inoculum was not acclimated. There was no
     information on controls and replicate test samples.
    Test temperature was 20 +/- 1 deg C.
F008 CLGETTS
F012 20
F020 119991
EOR
F002 27
F010 3.5
F004 11
F005 ME
F006 The Winkler method, which measures dissolved oxygen, is equivalent to the
     OECD 301D, Closed Bottle biodegradation test procedure.
F007 The Winkler method, which measures dissolved oxygen, is equivalent to the
     OECD 301D, Closed Bottle biodegradation test procedure.
F008 CLGETTS
F012 20
F020 119992
EOR
F002 27
F010 3.5
F004 11
F005 RE
F006 Lamb, C.B. and G.F. Jenkins. 1953. B.O.D. of Synthetic Organic Chemicals.
     Procedings of the 7th Industrial Waste Conference. pp. 326-339.
F007 Lamb, C.B. and G.F. Jenkins. 1953. B.O.D. of Synthetic Organic Chemicals.
     Procedings of the 7th Industrial Waste Conference. pp. 326-339.
F008 CLGETTS
F012 20
F020 119993
EOR
F002 27
F010 3.5
F004 11
F005 RL
F006 There is little information in the article describing how the study was
     conducted. This lack of information supports a reliability rating of 2.
     The results were comparable to reported data (Bridie, A.L., C.J.M. Wolff,
     and M. Winter. 1979. B
F007 There is little information in the article describing how the study was
     conducted. This lack of information supports a reliability rating of 2.
     The results were comparable to reported data (Bridie, A.L., C.J.M. Wolff,
     and M. Winter. 1979. BOD and COD of Some Petrochemicals. Water Research.
     13:627-630), which supports the acceptability of this study.
F008 CLGETTS
F012 20
F020 119994
EOR
F002 27
```

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F010 3.5
F004 11
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 119995
EOR
F002 27
F010 3.5
F004 11
F005 RS
F006 Day
            % Biodegradation (ThOD)
* *
* *
     5
                    0.0
* *
                    44.2
     10
* *
     15
                    69.2
* *
     20
                    72.3
* *
     30
                    73.2
* *
     40
                    75.4
* *
     50
                   77.0
F007 Day
           % Biodegradation (ThOD)
* *
* *
                    0.0
* *
    10
                    44.2
* *
                    69.2
    15
* *
     20
                    72.3
* *
     30
                    73.2
* *
     40
                   75.4
* *
                   77.0
     50
F008 CLGETTS
F012 20
F020 119996
EOR
F002 27
F010 3.5
F004 11
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 119997
EOR
F002 27
F010 3.5
F004 11
F005 TC
F006 An azide modification of the Winkler test method was used. The source of
     the inoculum from within the wastewater treatment plant was not stated.
     There was no information on controls and replicate test samples. Test
     material loading was 2.5
F007 An azide modification of the Winkler test method was used. The source of
     the inoculum from within the wastewater treatment plant was not stated.
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There was no information on controls and replicate test samples. Test
    material loading was 2.5 ppm.
    Test temperature was 20 deg C.
F008 CLGETTS
F012 20
F020 119998
EOR
F002 27
F010 3.6
F004 1
F005 RE
F006 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
     Reinhold Company, New York, NY, USA.
F007 Karel Verschueren. 1983. Handbook of Environmental Data on Organic
     Chemicals, 2nd Edition. Karel Verschueren (ed), pp. 301-302. Van Nostrand
     Reinhold Company, New York, NY, USA.
F008 HEDSET
F009 22-10-1992
F012 20
F020 119999
EOR
F002 27
F010 3.6
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120000
EOR
F002 27
F010 3.6
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120001
EOR
F002 27
F010 3.7
F004 1
F005 RE
F006 Bysshe, S.E. 1982. Biconcentration Factor in Aquatic Organisms. In:
    Handbook of Chemical Property Estimation Methods. Environmental Behavior
     of Organic Compounds. 5, 1-43. Lyman W.J., Reehl, W.F., and Rosenblatt,
    D.H. (eds). McGraw-Hill, Ne
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F007 Bysshe, S.E. 1982. Biconcentration Factor in Aquatic Organisms. In:
     Handbook of Chemical Property Estimation Methods. Environmental Behavior
     of Organic Compounds. 5, 1-43. Lyman W.J., Reehl, W.F., and Rosenblatt,
     D.H. (eds). McGraw-Hill, New York, NY, USA.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120002
EOR
F002 27
F010 3.7
F004 1
F005 RM
F006 A bioconcentration factor of 1.7 is calculated using a 2-butanol log
     octanol/water partition coefficient (Kow) of 0.61 as reported by Hansch
     et al., 1995 (Hansch, C., Leo, A., Hoekman, D. 1995. Exploring QSAR -
     Hydrophobic, Electronic, and
F007 A bioconcentration factor of 1.7 is calculated using a 2-butanol log
     octanol/water partition coefficient (Kow) of 0.61 as reported by Hansch
     et al., 1995 (Hansch, C., Leo, A., Hoekman, D. 1995. Exploring QSAR -
     Hydrophobic, Electronic, and Steric Constants. American Chemical Society,
     Washington, DC, USA.).
F008 HEDSET
F009 21-10-1992
F012 20
F020 120003
EOR
F002 27
F010 3.7
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120004
EOR
F002 27
F010 3.8
F004 1
F005 RE
F006 Thomas, R.G. 1982. Volatilization from Water. In: Handbook of Chemical
     Property Estimation Methods. Environmental Behavior of Organic Compounds.
     15, 1-34. Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds).
     McGraw-Hill, New York, NY, USA.
F007 Thomas, R.G. 1982. Volatilization from Water. In: Handbook of Chemical
     Property Estimation Methods. Environmental Behavior of Organic Compounds.
     15, 1-34. Lyman W.J., Reehl, W.F., and Rosenblatt, D.H. (eds).
     McGraw-Hill, New York, NY, USA.
F008 CLGETTS
F012 20
F020 120005
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EOR
F002 27
F010 3.8
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F012 20
F020 120006
EOR
F002 27
F010 4.1
F004 3
F005 RE
F006 Juhnke, I. and Luedemann, D. 1978. Results of the investigation of 200
     chemical compounds for acute fish
     toxicity with the golden orfe test. Z. Wasser. Abwasser
* *
    Forsch., 11(5):161-164.
F007 Juhnke, I. and Luedemann, D. 1978. Results of the investigation of 200
    chemical compounds for acute fish
* *
    toxicity with the golden orfe test. Z. Wasser. Abwasser
* *
    Forsch., 11(5):161-164.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120007
EOR
F002 27
F010 4.1
F004 3
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120008
EOR
F002 27
F010 4.1
F004 3
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
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F020 120009
EOR
F002 27
F010 4.1
F004 4
F005 ME
F006 Water used in the study came from Hammond Bay of Lake Huron and taken
     from a point source that was located 250 feet offshore at a depth of
     approximately nine feet. Test systems were 10L glass jars that contained
     5L of lake water. Test syste
F007 Water used in the study came from Hammond Bay of Lake Huron and taken
     from a point source that was located 250 feet offshore at a depth of
     approximately nine feet. Test systems were 10L glass jars that contained
     5L of lake water. Test systems were aerated. Up to six organisms were
     added to each test system. Larval lampreys (Petromyzon marinus) were
     collected by means of an electric shocker in the Ocqueoc River, Presque
     Isle County, Michigan, USA, and were held in running water in aquaria and
     samll "races" under conditions which simulated their natrual stream
    habitat.
* *
    Water pH ranged from 7.5 to 8.2; water temperature was 55+/-1 degrees F;
    dissolved oxygen ranged from 8.6 to 13.7 ppm; and free CO2 ranged from
     5.0 to 9.0 ppm. Treatment solutions used were 5.0, 1.0, and 0.1 ppm. A
     control system that only contained lake water was included. It was not
     stated whether the units were presented as a volume or weight.
F008 CLGETTS
F012 20
F020 120010
EOR
F002 27
F010 4.1
F004 4
F005 RE
F006 Applegate, V.C. et al. 1957. Toxicity of 4,346 chemicals to larval
     lampreys and fishes. Special Scientific Report--Fisheries No. 207. US
     Dept. of the Interior, US Fish and Wildlife Service, Washington, DC, USA.
F007 Applegate, V.C. et al. 1957. Toxicity of 4,346 chemicals to larval
     lampreys and fishes. Special Scientific Report--Fisheries No. 207. US
     Dept. of the Interior, US Fish and Wildlife Service, Washington, DC, USA.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120011
EOR
F002 27
F010 4.1
F004 4
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120012
EOR
F002 27
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F010 4.1
F004 4
F005 RS
F006 The highest treatment level tested was 5 ppm. No effects were observed at
     this level to laral lamprey.
F007 The highest treatment level tested was 5 ppm. No effects were observed at
     this level to laral lamprey.
F008 HEDSET
F012 20
F020 120013
EOR
F002 27
F010 4.1
F004 4
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120014
EOR
F002 27
F010 4.1
F004 5
F005 ME
F006 The test method used in this study is similar to the OECD 203 test
     guideline.
F007 The test method used in this study is similar to the OECD 203 test
     quideline.
F008 CLGETTS
F012 20
F020 120015
EOR
F002 27
F010 4.1
F004 5
F005 ME
F006 Trimmed Spearman-Karber Method (Hamilton, M.A., R.C. Russo, and R.V.
     Thurston. 1977. Trimmed Spearman-Karber Method for Estimating Median
     Lethal Concentrations in Toxicity Bioassays. Environ. Sci. Technol.
     11:714-719. Correction, 12:417, 19
F007 Trimmed Spearman-Karber Method (Hamilton, M.A., R.C. Russo, and R.V.
     Thurston. 1977. Trimmed Spearman-Karber Method for Estimating Median
     Lethal Concentrations in Toxicity Bioassays. Environ. Sci. Technol.
     11:714-719. Correction, 12:417, 1978.)
F008 CLGETTS
F012 20
F020 120016
EOR
F002 27
F010 4.1
F004 5
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F005 RE
F006 Geiger D.L. et al. 1986. Acute Toxicities of Organic Chemicals to Fathead
     Minnows (Pimephales promelas), Vol. 3. Center for Lake Superior
     Environmental Studies. University of Wisconsin-Superior, WS, USA.
F007 Geiger D.L. et al. 1986. Acute Toxicities of Organic Chemicals to Fathead
     Minnows (Pimephales promelas), Vol. 3. Center for Lake Superior
     Environmental Studies. University of Wisconsin-Superior, WS, USA.
F008 CLGETTS
F012 20
F020 120017
EOR
F002 27
F010 4.1
F004 5
F005 RM
F006 Purity:
              99%
F007 Purity: 99%
F008 CLGETTS
F012 20
F020 120018
EOR
F002 27
F010 4.1
F004 5
F005 RS
F006 96-hour LC50 = 3,670 \text{ mg/L} (95% CI 3,380 \text{ to } 3,990) based upon average,
     corrected measured values
* *
     Analytical method used was Gas-Liquid Chromatography
                            Fish Total
     Measured
* *
     Conc. (mg/L)
                        Mortality (@96 hrs)*
* *
       Control
F007 96-hour LC50 = 3,670 \text{ mg/L} (95% CI 3,380 \text{ to } 3,990) based upon average,
     corrected measured values
* *
     Analytical method used was Gas-Liquid Chromatography
* *
     Measured
                            Fish Total
* *
     Conc. (mg/L)
                        Mortality (@96 hrs)*
* *
       Control
                                 0
* *
                                 0
        1,018
* *
                                 0
        1,839
* *
                                 0
        2,630
* *
        4,258
                                16
* *
        6,667
                                2.0
* *
* *
     * 20 fish added at test initiation
F008 CLGETTS
F012 20
F020 120019
EOR
F002 27
F010 4.1
F004 5
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120020
```

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EOR
F002 27
F010 4.1
F004 5
F005 TC
F006 Treatment solutions were prepared by diluting a 3.7 g/L stock solution.
    Nominal sBA treatment levels were 1.05, 1.61, 2.48, 3.82, 5.88mg/L, which
    measured 1.02, 1.84, 2.63, 4.26 and 6.67mg/L, respectively.
     Control/dilution water was EPA Dul
F007 Treatment solutions were prepared by diluting a 3.7~\mathrm{g/L} stock solution.
    Nominal sBA treatment levels were 1.05, 1.61, 2.48, 3.82, 5.88mg/L, which
    measured 1.02, 1.84, 2.63, 4.26 and 6.67mg/L, respectively.
    Control/dilution water was EPA Duluth laboratory water. Twenty fish were
     tested per treatment and control. Tank volume = 2L. Control and treatment
     solution flow rate was equivalent to 18 chamber volumes per day.
    Mean test parameters were as follows: temperature = 24.4 deg. C;
    dissolved oxygen = 7.5 mg/L; pH = 7.8; fish age = 30 days; fish mean wt.
     = 0.09 g; fish mean length = 18.9 mm; fish loading = 0.0.90 g/L. Organism
     supplier was U.S. EPA Environmental Research Lab, Duluth, MN, USA.
F008 CLGETTS
F012 20
F020 120021
EOR
F002 27
F010 4.1
F004 6
F005 ME
F006 Interpolation from a graph of log concentration values versus percent
     mortality (APHA, 1971).
F007 Interpolation from a graph of log concentration values versus percent
     mortality (APHA, 1971).
F008 CLGETTS
F012 20
F020 120022
EOR
F002 27
F010 4.1
F004 6
F005 RE
F006 Bridie, A.L., C.J.M. Wolff, and M. Winter. 1979. The Acute Toxicity of
     Some Petrochemicals to Goldfish. Water Research. 13:623-626.
F007 Bridie, A.L., C.J.M. Wolff, and M. Winter. 1979. The Acute Toxicity of
     Some Petrochemicals to Goldfish. Water Research. 13:623-626.
F008 CLGETTS
F012 20
F020 120023
EOR
F002 27
F010 4.1
F004 6
F005 RL
F006 It is unclear from the article as to whether the reported result is based
     on nominal or measured values. This lack of information, in conjunction
     with the absence of selected test parameters and results (i.e,, exposure
     concentrations, morta
F007 It is unclear from the article as to whether the reported result is based
     on nominal or measured values. This lack of information, in conjunction
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with the absence of selected test parameters and results (i.e,, exposure
     concentrations, mortality), supports a reliability rating of 2. The
    reported value in this study is comparable to a reported fathead minnow
     acute value for sBA: 24-hour LC50 = ~4,000 mg/L (Geiger D.L. et al. 1986.
    Acute Toxicities of Organic Chemicals to Fathead Minnows (Pimephales
    promelas), Vol. 3. Center for Lake Superior Environmental Studies.
     University of Wisconsin-Superior, WS, USA.).
F008 CLGETTS
F012 20
F020 120024
EOR
F002 27
F010 4.1
F004 6
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 120025
EOR
F002 27
F010 4.1
F004 6
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120026
EOR
F002 27
F010 4.1
F004 6
F005 TC
F006 A series of treatment solutions were prepared, but their concentrations
    were not reported. Ten fish were tested per treatment. Tank volume = 25L.
    Treatment solutions were aerated during the test. Treatment
     concentrations were analytically v
F007 A series of treatment solutions were prepared, but their concentrations
     were not reported. Ten fish were tested per treatment. Tank volume = 25L.
     Treatment solutions were aerated during the test. Treatment
     concentrations were analytically verified at test start and termination,
    but the method of analysis was not reported. Reported results were not
    distinguished as being based on nominal or measured values and there is
    no control information.
    Test parameters were as follows: temperature = 20+/-1 deg. C; dissolved
     oxygen did not fall below 4 mg/L; initial pH = 7.0; fish average wt. =
     3.3+/-1.0 g; fish average length = 6.2+/-0.7 cm.
F008 CLGETTS
F012 20
F020 120027
EOR
F002 27
F010 4.1
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F004 7
F005 RE
F006 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F007 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120028
EOR
F002 27
F010 4.1
F004 7
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 CLGETTS
F012 20
F020 120029
EOR
F002 27
F010 4.1
F004 7
F005 RM
F006 Test Type: Acute Fish Toxicity Calculation
F007 Test Type: Acute Fish Toxicity Calculation
F008 CLGETTS
F012 20
F020 120030
EOR
F002 27
F010 4.1
F004 7
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., NJ, USA
F007 ExxonMobil Biomedical Sciences, Inc., NJ, USA
F008 CLGETTS
F012 20
F020 120031
EOR
F002 27
F010 4.1
F004 7
F005 TC
F006 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used
F007 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
    Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
    Entry System) structure used with the model was: OC(CC)C. Additional
    data calculated by the model include molecular weight , 74.12, and water
     solubility, 8,625 mg/L (@25C).
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The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.

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Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American
     Chemical Society. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120032
EOR
F002 27
F010 4.2
F004 1
F005 RE
F006 Bringmann, G. and Kuhn, R. 1977. Results of the damaging
     effect of water pollutants on Daphnia magna. Z. Wasser
* *
     Abwasser Forsch., 10(5):161-166.
F007 Bringmann, G. and Kuhn, R. 1977. Results of the damaging
     effect of water pollutants on Daphnia magna. Z. Wasser
     Abwasser Forsch., 10(5):161-166.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120033
EOR
F002 27
F010 4.2
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120034
EOR
F002 27
F010 4.2
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120035
EOR
F002 27
F010 4.2
F004 2
F005 ME
F006 None applied. The EC50 was calculated arithmetically from the
     concentration/effect ratio.
F007 None applied. The EC50 was calculated arithmetically from the
     concentration/effect ratio.
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F008 CLGETTS

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F012 20
F020 120036
EOR
F002 27
F010 4.2
F004 2
F005 ME
F006 The test method used in this study is similar to the OECD 202 test
     quideline.
F007 The test method used in this study is similar to the OECD 202 test
     guideline.
F008 CLGETTS
F012 20
F020 120037
EOR
F002 27
F010 4.2
F004 2
F005 RE
F006 Kuhn, R., M. Pattard, K-D. Pernak and A. Winter. 1989. Results of the
     Harmful Effects of Selected Water Pollutants (Anilines, Phenols,
     Aliphatic Compounds) to Daphnia magna. Water Research. 14:495-499.
F007 Kuhn, R., M. Pattard, K-D. Pernak and A. Winter. 1989. Results of the
     Harmful Effects of Selected Water Pollutants (Anilines, Phenols,
     Aliphatic Compounds) to Daphnia magna. Water Research. 14:495-499.
F008 CLGETTS
F012 20
F020 120038
EOR
F002 27
F010 4.2
F004 2
F005 RL
F006 Although the method was described in the article, data were not provided
     on the test parameters or results from individual treatment and control
     solutions. This lack of information supports a reliability rating of 2.
F007 Although the method was described in the article, data were not provided
     on the test parameters or results from individual treatment and control
     solutions. This lack of information supports a reliability rating of 2.
F008 CLGETTS
F012 20
F020 120039
EOR
F002 27
F010 4.2
F004 2
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 120040
EOR
F002 27
F010 4.2
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F004 2
F005 RS
F006 48-hour EC50 = 4,227 mg/L (95% CI 1,859 to 7,143) based on nominal values
     The authors considered the test valid because fewer than 10% of the
     organisms in the control were immobile, the pH value was not below 7.0,
     and the DO was not below 4
F007 48-hour EC50 = 4,227 mg/L (95% CI 1,859 to 7,143) based on nominal values
     The authors considered the test valid because fewer than 10% of the
     organisms in the control were immobile, the pH value was not below 7.0,
     and the DO was not below 4.0 mg/L at test termination.
F008 CLGETTS
F012 20
F020 120041
EOR
F002 27
F010 4.2
F004 2
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120042
EOR
F002 27
F010 4.2
F004 2
F005 TC
F006 The test procedure used daphnids 6 to 24 hours old. Organisms were not
     fed during the test. The treatment medium was defined and had a total
     hardness of 2.4 mmol/L, a calcium to magnesium ratio of 4:1, a sodium to
     potassium ratio of 10:1, a
F007 The test procedure used daphnids 6 to 24 hours old. Organisms were not
     fed during the test. The treatment medium was defined and had a total
    hardness of 2.4 mmol/L, a calcium to magnesium ratio of 4:1, a sodium to
    potassium ratio of 10:1, and an initial pH value of 8.0+/-0.2. The test
    solution temperature was 20°C.
    The test systems used 50 ml beakers. Treatment solutions and controls
    were prepared in duplicate. The organism loading rate was no less than
     one animal per 2 ml test medium. 20 organisms were tested per treatment
     and control. Treatment solutions were prepared from a stock sBA solution.
    Dissolved oxygen (DO) and pH were measured at test termination although
     specific values were not supplied for this test.
F008 CLGETTS
F012 20
F020 120043
EOR
F002 27
F010 4.2
F004 3
F005 ME
F006 Probit analysis procedure using Statistical Analysis System (SAS)
     software (SAS Institute, 1989).
F007 Probit analysis procedure using Statistical Analysis System (SAS)
     software (SAS Institute, 1989).
F008 CLGETTS
F012 20
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F020 120044
EOR
F002 27
F010 4.2
F004 3
F005 RE
F006 Schultz, W. and M. Tichy. 1993. Structure-Toxicity Relationships for
     Unsaturated Alcohols to Tetrahymena pyriformis: C5 and C6 Analogs and
     Primary Propargylic Alcohols. Environ. Contam. and Toxicology. 51:681-688.
F007 Schultz, W. and M. Tichy. 1993. Structure-Toxicity Relationships for
    Unsaturated Alcohols to Tetrahymena pyriformis: C5 and C6 Analogs and
     Primary Propargylic Alcohols. Environ. Contam. and Toxicology. 51:681-688.
F008 CLGETTS
F012 20
F020 120045
EOR
F002 27
F010 4.2
F004 3
F005 RL
F006 A non-standardized method was referenced in the article. Data were not
    provided on the test parameters or results from individual treatment and
     control solutions. This lack of information supports a reliability rating
     of 2.
F007 A non-standardized method was referenced in the article. Data were not
    provided on the test parameters or results from individual treatment and
     control solutions. This lack of information supports a reliability rating
     of 2.
F008 CLGETTS
F012 20
F020 120046
EOR
F002 27
F010 4.2
F004 3
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 120047
EOR
F002 27
F010 4.2
F004 3
F006 48-hour EC50 for growth = 3,196 mg/L based on nominal values
     Population density (growth) was measured spectrophotometrically as
     absorbance at 540 nm.
F007 48-hour EC50 for growth = 3,196 mg/L based on nominal values
     Population density (growth) was measured spectrophotometrically as
     absorbance at 540 nm.
F008 CLGETTS
F012 20
F020 120048
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EOR
F002 27
F010 4.2
F004 3
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120049
EOR
F002 27
F010 4.2
F004 3
F005 TC
F006 Axenic cultures of Tetrahymena pyrifomis were used in the test. Only
     48-hour population densities were measured. Treatment solutions and
     controls were prepared in duplicate. Treatment solutions were prepared
     from a stock sBA solution.
F007 Axenic cultures of Tetrahymena pyrifomis were used in the test. Only
     48-hour population densities were measured. Treatment solutions and
     controls were prepared in duplicate. Treatment solutions were prepared
     from a stock sBA solution.
F008 CLGETTS
F012 20
F020 120050
EOR
F002 27
F010 4.2
F004 4
F005 RE
F006 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F007 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120051
EOR
F002 27
F010 4.2
F004 4
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 CLGETTS
F012 20
F020 120052
EOR
F002 27
F010 4.2
F004 4
F005 RM
F006 Test Type: Acute Daphnid Toxicity Calculation
F007 Test Type: Acute Daphnid Toxicity Calculation
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F008 CLGETTS
F012 20
F020 120053
EOR
F002 27
F010 4.2
F004 4
F005 S0
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120054
EOR
F002 27
F010 4.2
F004 4
F005 TC
F006 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used
F007 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
    Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
    Entry System) structure used with the model was: OC(CC)C. Additional
    data calculated by the model include molecular weight , 74.12, and water
     solubility, 8,625 \text{ mg/L} (@25C).
    The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.
     Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American
     Chemical Society. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120055
EOR
F002 27
F010 4.3
F004 2
F005 RE
F006 Bringmann, G. and Kuhn, R. 1976. Vergleichende Befunde der Schadwirkung
     wassergefahrdender Stoffe gegen Bakterien (Pseudomonas putida) und
     Blaualgen (Microcystis aeruginosa). Gwf-Wasser/Abwasser, Vol. 117.
F007 Bringmann, G. and Kuhn, R. 1976. Vergleichende Befunde der Schadwirkung
     wassergefahrdender Stoffe gegen Bakterien (Pseudomonas putida) und
     Blaualgen (Microcystis aeruginosa). Gwf-Wasser/Abwasser, Vol. 117.
F008 HEDSET
F009 16-02-1994
F012 20
F020 120056
EOR
F002 27
F010 4.3
F004 2
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
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commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120057
EOR
F002 27
F010 4.3
F004 2
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 HEDSET
F012 20
F020 120058
EOR
F002 27
F010 4.3
F004 3
F005 RE
F006 Jones, H.R. 1971. Environmental control in the organic and petrochemical
     industries. Noyes Data Corporation.
F007 Jones, H.R. 1971. Environmental control in the organic and petrochemical
     industries. Noyes Data Corporation.
F008 HEDSET
F009 31-03-1994
F012 20
F020 120059
F:OR
F002 27
F010 4.3
F004 3
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120060
EOR
F002 27
F010 4.3
F004 3
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 HEDSET
F012 20
F020 120061
EOR
F002 27
F010 4.3
F004 4
F005 ME
F006 None applied. The toxicity threshold (TT) was determined graphically by
     plotting the highest non-toxic concentration versus its mean extinction
     value against the lowest toxic concentration versus its mean extinction
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value and calculating th
F007 None applied. The toxicity threshold (TT) was determined graphically by
     plotting the highest non-toxic concentration versus its mean extinction
    value against the lowest toxic concentration versus its mean extinction
    value and calculating the toxicant concentration at 3% below the no
     effect level.
F008 HEDSET
F012 20
F020 120062
EOR
F002 27
F010 4.3
F004 4
F005 RE
F006 Bringmann, G. and R. Kuhn. 1980. Comparison of the Toxicity Thresholds of
     Water Pollutants to Bacteria, Algae, and Protozoa in the Cell
     Multiplication Inhibition Test. Water Research. 14:231-241.
F007 Bringmann, G. and R. Kuhn. 1980. Comparison of the Toxicity Thresholds of
     Water Pollutants to Bacteria, Algae, and Protozoa in the Cell
     Multiplication Inhibition Test. Water Research. 14:231-241.
F008 HEDSET
F012 20
F020 120063
EOR
F002 27
F010 4.3
F004 4
F005 RL
F006 Although a non-standardized method was described in the article, data
     were not provided on the test parameters, replication, or results from
     individual treatment and control solutions. This lack of information
     supports a reliability rating
F007 Although a non-standardized method was described in the article, data
     were not provided on the test parameters, replication, or results from
     individual treatment and control solutions. This lack of information
     supports a reliability rating of 2.
F008 HEDSET
F012 20
F020 120064
EOR
F002 27
F010 4.3
F004 4
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120065
EOR
F002 27
F010 4.3
F004 4
F005 RM
F006 Test Type: Static Toxicity Test
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F007 Test Type: Static Toxicity Test
F008 HEDSET
F012 20
F020 120066
EOR
F002 27
F010 4.3
F004 4
F005 RS
F006 7-day TT (toxicity threshold) for growth = 95 mg/L based on nominal
     values.
    The TT value for growth is calculated by identifying the treatment level
     that is greater or equal to 3% below the treatment level that did not
     exhibit toxic effects
F007 7-day TT (toxicity threshold) for growth = 95 mg/L based on nominal
     values.
     The TT value for growth is calculated by identifying the treatment level
     that is greater or equal to 3% below the treatment level that did not
     exhibit toxic effects as measured by the extinction of primary light of
     monochromatic radiation at 578 nm.
F008 HEDSET
F012 20
F020 120067
EOR
F002 27
F010 4.3
F004 4
F005 S0
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 HEDSET
F012 20
F020 120068
EOR
F002 27
F010 4.3
F004 4
F005 TC
F006 Treatment solutions were prepared by diluting a stock sBA solution.
     Testing was conducted in metal capped, 300 ml Erlenmeyer flasks
     containing 50 ml of treatment solution. Treatment solutions contained
     sBA, cells, double distilled water, a
F007 Treatment solutions were prepared by diluting a stock sBA solution.
     Testing was conducted in metal capped, 300 ml Erlenmeyer flasks
     containing 50 ml of treatment solution. Treatment solutions contained
     sBA, cells, double distilled water, and a sterile, defined nutrient
    medium. The control solution contained nutrient medium, to which sterile
     double distilled water was added. Growth inhibition measurements were
     only determined on day 7.
    Cell growth was determined by using a turbidimetric procedure that
     measured primary light extinction (monochromatic radiation at 578 nm)
     through a cell suspension of 10 mm thickness.
F008 HEDSET
F012 20
F020 120069
EOR
F002 27
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F010 4.3
F004 5
F005 RE
F006 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F007 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F008 HEDSET
F012 20
F020 120070
EOR
F002 27
F010 4.3
F004 5
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 HEDSET
F012 20
F020 120071
EOR
F002 27
F010 4.3
F004 5
F005 RM
F006 Test Type: Green Alga Toxicity Calculation
F007 Test Type: Green Alga Toxicity Calculation
F008 HEDSET
F012 20
F020 120072
EOR
F002 27
F010 4.3
F004 5
F005 RS
F006 96-hour EC50 = 625 mg/L
F007 96-hour EC50 = 625 mg/L
F008 HEDSET
F012 20
F020 120073
EOR
F002 27
F010 4.3
F004 5
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., NJ, USA
F007 ExxonMobil Biomedical Sciences, Inc., NJ, USA
F008 HEDSET
F012 20
F020 120074
EOR
F002 27
F010 4.3
F004 5
F005 TC
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F006 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used
F007 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used with the model was: OC(CC)C. Additional
     data calculated by the model include molecular weight , 74.12, and water
     solubility, 8,625 mg/L (@25C).
     The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.
     Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American
     Chemical Society. Washington, DC, USA.
F008 HEDSET
F012 20
F020 120075
EOR
F002 27
F010 4.3
F004 6
F005 RE
F006 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F007 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F008 HEDSET
F012 20
F020 120076
EOR
F002 27
F010 4.3
F004 6
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 HEDSET
F012 20
F020 120077
EOR
F002 27
F010 4.3
F004 6
F005 RS
F006 *Chronic value
F007 *Chronic value
F008 HEDSET
F012 20
F020 120078
EOR
F002 27
F010 4.3
F004 6
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
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F008 HEDSET
F012 20
F020 120079
EOR
F002 27
F010 4.3
F004 6
F005 TC
F006 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used
F007 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used with the model was: OC(CC)C. Additional
     data calculated by the model include molecular weight , 74.12, and water
     solubility, 8,625 mg/L (@25C).
    The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.
     Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American
     Chemical Society. Washington, DC, USA.
F008 HEDSET
F012 20
F020 120080
EOR
F002 27
F010 4.4
F004 1
F005 RE
F006 Bringmann, G. and R. Kuhn. 1980. Comparison of the toxicity thresholds
     of water pollutants to bacteria, algae and protozoa in the cell
     multiplication inhibition test. Water Research, 14:231-241.
F007 Bringmann, G. and R. Kuhn. 1980. Comparison of the toxicity thresholds
     of water pollutants to bacteria, algae and protozoa in the cell
    multiplication inhibition test. Water Research, 14:231-241.
F008 HEDSET
F009 20-10-1992
F012 20
F020 120081
EOR
F002 27
F010 4.4
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120082
EOR
F002 27
F010 4.4
F004 1
F005 S0
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
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F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 HEDSET
F012 20
F020 120083
EOR
F002 27
F010 4.4
F004 2
F005 RE
F006 Environmental Health Criteria 65: Butanols: four isomers:
     1-butanol, 2-butanol, tert-butanol, isobutanol. 1987. IPCS
* *
     International Programme on Chemical Safety, World Health
* *
     Organization.
F007 Environmental Health Criteria 65: Butanols: four isomers:
     1-butanol, 2-butanol, tert-butanol, isobutanol. 1987. IPCS
* *
     International Programme on Chemical Safety, World Health
* *
     Organization.
F008 HEDSET
F009 20-10-1992
F012 20
F020 120084
EOR
F002 27
F010 4.4
F004 2
F005 S0
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120085
EOR
F002 27
F010 4.4
F004 3
F005 RE
F006 Bringmann, G. and Kuhn, R. 1980. Bestimmung der biologischen Schadwirkung
     wassergefahrdender Stoffe gegen Protozoen. II. Bakterienfressende
     Ciliaten. Z. Wasser/Abwasser Forsch., 1:26-31.
F007 Bringmann, G. and Kuhn, R. 1980. Bestimmung der biologischen Schadwirkung
     wassergefahrdender Stoffe gegen Protozoen. II. Bakterienfressende
     Ciliaten. Z. Wasser/Abwasser Forsch., 1:26-31.
F008 CLGETTS
F012 20
F020 120086
EOR
F002 27
F010 4.4
F004 3
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
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F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 120087
EOR
F002 27
F010 4.4
F004 3
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120088
EOR
F002 27
F010 4.4
F004 4
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 120089
EOR
F002 27
F010 4.4
F004 4
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120090
EOR
F002 27
F010 4.4
F004 5
F005 RE
F006 Bringmann, G. and Kuhn, R. 1980. Comparison of the toxicity thresholds of
     water pollutants to bacteria, algae and protozoa in the cell
     multiplication inhibition test. Water Research, 14:231-241.
F007 Bringmann, G. and Kuhn, R. 1980. Comparison of the toxicity thresholds of
     water pollutants to bacteria, algae and protozoa in the cell
     multiplication inhibition test. Water Research, 14:231-241.
F008 CLGETTS
F012 20
F020 120091
EOR
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F002 27
F010 4.4
F004 5
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
    commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 120092
EOR
F002 27
F010 4.4
F004 5
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120093
EOR
F002 27
F010 4.5.1
F004 1
F005 RE
F006 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F007 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120094
EOR
F002 27
F010 4.5.1
F004 1
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 CLGETTS
F012 20
F020 120095
EOR
F002 27
F010 4.5.1
F004 1
F005 RM
F006 Test Type: Chronic Fish Toxicity Calculation
    * Chronic Value
F007 Test Type: Chronic Fish Toxicity Calculation
** * Chronic Value
F008 CLGETTS
F012 20
F020 120096
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EOR
F002 27
F010 4.5.1
F004 1
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120097
EOR
F002 27
F010 4.5.1
F004 1
F005 TC
F006 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used
F007 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
    Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
    Entry System) structure used with the model was: OC(CC)C. Additional
    data calculated by the model include molecular weight , 74.12, and water
     solubility, 8,625 mg/L (@25C).
    The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.
    Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American
     Chemical Society. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120098
EOR
F002 27
F010 4.5.2
F004 1
F005 RE
F006 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
    Assessment Division. Washington, DC, USA.
F007 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120099
EOR
F002 27
F010 4.5.2
F004 1
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 CLGETTS
F012 20
F020 120100
EOR
F002 27
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F010 4.5.2
F004 1
F005 RM
F006 Test Type: Chronic Daphnid Toxicity Calculation
F007 Test Type: Chronic Daphnid Toxicity Calculation
F008 CLGETTS
F012 20
F020 120101
EOR
F002 27
F010 4.5.2
F004 1
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120102
EOR
F002 27
F010 4.5.2
F004 1
F005 TC
F006 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used
F007 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used with the model was: OC(CC)C. Additional
     data calculated by the model include molecular weight , 74.12, and water
     solubility, 8,625 mg/L (@25C).
* *
     The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.
     Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American
     Chemical Society. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120103
EOR
F002 27
F010 4.6.2
F004 1
F005 RE
F006 World Health Organization. 1987. Environmental Health Criteria 65.
F007 World Health Organization. 1987. Environmental Health Criteria 65.
F008 HEDSET
F009 20-10-1992
F012 20
F020 120109
EOR
F002 27
F010 4.6.2
F004 1
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
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F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120110
EOR
F002 27
F010 4.6.2
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120111
EOR
F002 27
F010 4.6.2
F004 2
F005 RE
F006 World Health Organization. 1987. Environmental Health Criteria 65.
F007 World Health Organization. 1987. Environmental Health Criteria 65.
F008 HEDSET
F009 20-10-1992
F012 20
F020 120112
EOR
F002 27
F010 4.6.2
F004 2
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120113
EOR
F002 27
F010 4.6.2
F004 2
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
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     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
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F020 120114
EOR
F002 27
F010 4.6.2
F004 3
F005 ME
F006 Stage at application: seedling, on intact plant; addition to
     growth meduim, in culture flask, dose = 1%.
F007 Stage at application: seedling, on intact plant; addition to
     growth meduim, in culture flask, dose = 1%.
F008 CLGETTS
F012 20
F020 120115
EOR
F002 27
F010 4.6.2
F004 3
F005 RE
F006 Macht, D.I. and Meyer, J.D. 1933. Am. J. Botany, 20:145-149.
F007 Macht, D.I. and Meyer, J.D. 1933. Am. J. Botany, 20:145-149.
F008 HEDSET
F009 15-02-1994
F012 20
F020 120116
EOR
F002 27
F010 4.6.2
F004 3
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120117
EOR
F002 27
F010 4.6.2
F004 3
F005 RS
F006 73% size decrease of root
F007 73% size decrease of root
F008 HEDSET
F012 20
F020 120118
EOR
F002 27
F010 4.6.2
F004 3
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
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     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
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F008 CLGETTS
F009 09-01-2002
F012 20
F020 120119
EOR
F002 27
F010 4.6.2
F004 4
F005 ME
F006 stage at application: seedling, on intact plant, in
    environmental chamber, application by fumigation: 2 cm3/1.
F007 stage at application: seedling, on intact plant, in
    environmental chamber, application by fumigation: 2 cm3/1.
F008 CLGETTS
F012 20
F020 120120
EOR
F002 27
F010 4.6.2
F004 4
F005 RE
F006 Miller, L.P. 1934. Contr. Boy. T., 6:279-296.
F007 Miller, L.P. 1934. Contr. Boy. T., 6:279-296.
F008 HEDSET
F009 15-02-1994
F012 20
F020 120121
EOR
F002 27
F010 4.6.2
F004 4
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120122
EOR
F002 27
F010 4.6.2
F004 4
F005 RS
F006 20% respiration increase, on tuber
F007 20% respiration increase, on tuber
F008 HEDSET
F012 20
F020 120123
EOR
F002 27
F010 4.6.2
F004 4
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
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F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120124
EOR
F002 27
F010 4.6.2
F004 5
F005 ME
F006 Stage at application: seedling, on excised organ; addition
** to growth medium, dose: 5 \times 10 E-2 M.
F007 Stage at application: seedling, on excised organ; addition
** to growth medium, dose: 5 x 10 E-2 M.
F008 CLGETTS
F012 20
F020 120125
EOR
F002 27
F010 4.6.2
F004 5
F005 RE
F006 Gudjonsdottir, S. and Burstrom, H. 1962. Physl. Plant, 15:498-504.
F007 Gudjonsdottir, S. and Burstrom, H. 1962. Physl. Plant, 15:498-504.
F008 HEDSET
F009 15-02-1994
F012 20
F020 120126
EOR
F002 27
F010 4.6.2
F004 5
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120127
EOR
F002 27
F010 4.6.2
F004 5
F005 RS
F006 50% size decrease of cells
F007 50% size decrease of cells
F008 HEDSET
F012 20
F020 120128
EOR
F002 27
F010 4.6.2
F004 5
F005 SO
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F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120129
EOR
F002 27
F010 4.6.2
F004 6
F005 ME
F006 Stage at application: seedling, on excised organ; addition
** to growth medium, dose: 4 x 10E-2 M
F007 Stage at application: seedling, on excised organ; addition
     to growth medium, dose: 4 x 10E-2 M
F008 CLGETTS
F012 20
F020 120130
EOR
F002 27
F010 4.6.2
F004 6
F005 RE
F006 Gudjonsdottir, S. and Burstrom, H. Physl. Plant, 15:498-504.
F007 Gudjonsdottir, S. and Burstrom, H. Physl. Plant, 15:498-504.
F008 HEDSET
F009 15-02-1994
F012 20
F020 120131
EOR
F002 27
F010 4.6.2
F004 6
F005 RM
F006 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 HEDSET
F012 20
F020 120132
EOR
F002 27
F010 4.6.2
F004 6
F005 RS
F006 50% decrease in number of cells
F007 50% decrease in number of cells
F008 HEDSET
F012 20
F020 120133
EOR
F002 27
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F010 4.6.2
F004 6
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
* *
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120134
EOR
F002 27
F010 4.6.3
F004 1
F005 RE
F006 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F007 Cash, G. and V. Nabholz.. 1999. ECOSAR Classes for Microsoft Windows,
     ECOWIN v0.99e. U.S. Environmental Protection Agency, OPPT - Risk
     Assessment Division. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120104
F:OR
F002 27
F010 4.6.3
F004 1
F005 RL
F006 Rated 2 for reliability due to calculated value.
F007 Rated 2 for reliability due to calculated value.
F008 CLGETTS
F012 20
F020 120105
EOR
F002 27
F010 4.6.3
F004 1
F005 RM
F006 Test Type: Earthworm Toxicity Calculation
F007 Test Type: Earthworm Toxicity Calculation
F008 CLGETTS
F012 20
F020 120106
EOR
F002 27
F010 4.6.3
F004 1
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120107
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EOR
F002 27
F010 4.6.3
F004 1
F005 TC
F006 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
     Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used
F007 A log Kow (octanol/water partition coefficient) value and a chemical
     structure are needed to complete the calculation with this model. The log
    Kow value used was 0.61. The SMILES (Simplified Molecular Input Line
     Entry System) structure used with the model was: OC(CC)C. Additional
     data calculated by the model include molecular weight , 74.12, and water
     solubility, 8,625 mg/L (@25C).
    The reference for log Kow is: Hansch, C., A. Leo, and D. Hoekman. 1995.
     Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. American
     Chemical Society. Washington, DC, USA.
F008 CLGETTS
F012 20
F020 120108
EOR
F002 27
F010 4.6.4
F004 2
F005 ME
F006 Median lethal concentration was calculated as a projection from the least
     square linear regression on log transformed nominal concentration data
     and probit transformed percent effect data.
F007 Median lethal concentration was calculated as a projection from the least
     square linear regression on log transformed nominal concentration data
     and probit transformed percent effect data.
F008 CLGETTS
F012 20
F020 120135
EOR
F002 27
F010 4.6.4
F004 2
F005 RE
F006 de Zwart, D. and Slooff, W. 1987. Toxicity of Mixtures of Heavy Metals
     and Petrochemicals to Xenopus laevis. Bull. Environ. Contam. Toxicol.,
     38:345-351.
F007 de Zwart, D. and Slooff, W. 1987. Toxicity of Mixtures of Heavy Metals
     and Petrochemicals to Xenopus laevis. Bull. Environ. Contam. Toxicol.,
     38:345-351.
F008 CLGETTS
F012 20
F020 120136
EOR
F002 27
F010 4.6.4
F004 2
F005 RL
F006 The test procedure was generally described in the article. Data were not
     provided on the test parameters or results from individual treatment and
     control solutions. This lack of information supports a reliability rating
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of 2.
F007 The test procedure was generally described in the article. Data were not
     provided on the test parameters or results from individual treatment and
     control solutions. This lack of information supports a reliability rating
     of 2.
F008 CLGETTS
F012 20
F020 120137
EOR
F002 27
F010 4.6.4
F004 2
F005 RM
F006 Test Type: Static Acute Toxicity Test
    Analytical Monitoring: No
* *
    Results based on nominal treatment values.
     Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F007 Test Type: Static Acute Toxicity Test
* *
    Analytical Monitoring: No
* *
    Results based on nominal treatment values.
    Information on sBA purity is not available; assumed to have used a
     commercial grade of >/= 99%.
F008 CLGETTS
F012 20
F020 120138
EOR
F002 27
F010 4.6.4
F004 2
F005 SO
F006 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F007 ExxonMobil Biomedical Sciences, Inc., New Jersey, USA
F008 CLGETTS
F012 20
F020 120139
EOR
F002 27
F010 4.6.4
F004 2
F005 TC
F006 The test procedure used 3 to 4 week old larvae. The organisms were
     exposed in covered all-glass aquaria containing 1L of Dutch Standard
     Water at a temperature of 20+/-1C. The test material was added once at
     the beginning of the study. Five
F007 The test procedure used 3 to 4 week old larvae. The organisms were
     exposed in covered all-glass aquaria containing 1L of Dutch Standard
     Water at a temperature of 20+/-1C. The test material was added once at
     the beginning of the study. Five concentration levels were evaluated with
     10 organisms per level. The 5 levels had a factorial difference of 1.5.
    Mortality was only recorded after 48 hours.
    The test species, Xenopus laevis, was identified in the article as a
    clawed toad. However, it is correctly identified as a clawed frog in this
    robust summary.
F008 CLGETTS
F012 20
F020 120140
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EOR
F002 27
F010 5.0
F004 1
F005 CL
F006 In the development of the model, the authors confirmed that a limited
     amount of sBA could be recovered as glucuronide in the urine. Most of the
     alcohol appears to undergo oxidation via alcohol dehydrogenase to its
     corresponding ketone, MEK.
F007 In the development of the model, the authors confirmed that a limited
     amount of sBA could be recovered as glucuronide in the urine. Most of the
     alcohol appears to undergo oxidation via alcohol dehydrogenase to its
     corresponding ketone, MEK. The results showed a limited amount of the
    ketone undergoes a backward reduction to its parent alcohol, sBA.
     and 2,3-BD are found to be common metabolites of sBA and MEK. The model
    was able to simulate blood concentrations and elimination of all 4
     compounds after oral administration of sBA, and the results after i.v. of
     3H-2B and 2,3-BD. AUC analysis suggested that the quantities of 3H-2B
    and 2,3-BD formed from oral doses of sBA and MEK are comparable. The
    results supported the estimation in that no significant difference in the
    AUC of MEK was observed after dosing with either 1776 mg/kg of sBA or
     1690 mg/kg of MEK (10,899 \pm 824 vs. 9868 \pm 566 mg-hr /liter,
     respectively).
F020 255640
EOR
F002 27
F010 5.0
F004 1
F005 ME
F006 Statistical Methods
     Student's t-test was used for statistical evaluation of differences
    between two means. In the pharmacokinetic model, the differential
     equations were solved numerically by a Hammings Predictor-Correction
    method with a Ru
F007 Statistical Methods
     Student's t-test was used for statistical evaluation of differences
     between two means. In the pharmacokinetic model, the differential
     equations were solved numerically by a Hammings Predictor-Correction
     method with a Runge-Kutta starter.
F020 255637
EOR
F002 27
F010 5.0
F004 1
F005 RE
F006 Dietz FK, Rodrigues-Giaxola M, Traiger GJ, Stella VJ and Himmelstein KJ
     (1981). Pharmacokinetics of 2-Butanol and its metabolites in the rat.
     Journal of Pharmacokinetics and Biopharmaceutics 9(5), 553-576.
F007 Dietz FK, Rodrigues-Giaxola M, Traiger GJ, Stella VJ and Himmelstein KJ
     (1981). Pharmacokinetics of 2-Butanol and its metabolites in the rat.
     Journal of Pharmacokinetics and Biopharmaceutics 9(5), 553-576.
F020 255642
EOR
F002 27
F010 5.0
F004 1
F005 RL
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F006 No circumstances occurred that would have affected the quality or
     integrity of the data. Test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
F007 No circumstances occurred that would have affected the quality or
     integrity of the data. Test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
F020 255641
EOR
F002 27
F010 5.0
F004 1
F005 RM
F006 Dietz et al. (1981) selected doses for MEK and sBA based on the earlier
    pharmacokinetic analysis (Traiger and Bruckner, 1976) that established
     approximately 96% of an administered dose of sBA was oxidized in vivo to
     MEK. Thus 1960 mg/kg (2.
F007 Dietz et al. (1981) selected doses for MEK and sBA based on the earlier
     pharmacokinetic analysis (Traiger and Bruckner, 1976) that established
     approximately 96% of an administered dose of sBA was oxidized in vivo to
    MEK. Thus 1960 mg/kg (2.1 ml/kg) of MEK estimates the amount of MEK
     formed in vivo from a dose of 1776 mg/kg (2.2 ml/kg) of sBA. Male rats
     were given oral doses of sBA, MEK, 2-3 B-D (or i.v.), or 3H-2B (i.v.) and
     serial blood samples were collected for up to 30 hours following
     treatment.
F020 255639
EOR
F002 27
F010 5.0
F004 1
F005 RM
F006 Test Type: Metabolism and Pharmacokinetic Evaluation
     Strain: Sprague-Dawley
     Dose Groups/Concentrations: sBA: 2.2 ml/kg or 1776 mg/kg as a 22%
     aqueous solution (oral). MEK: 2.1ml/kg or 1690 mg/kg as a 21% aqueous
     solution (oral). 2,3-BD
F007 Test Type: Metabolism and Pharmacokinetic Evaluation
* *
    Strain: Spraque-Dawley
* *
    Dose Groups/Concentrations: sBA: 2.2 ml/kg or 1776 mg/kg as a 22%
    aqueous solution (oral). MEK: 2.1ml/kg or 1690 mg/kg as a 21% aqueous
     solution (oral). 2,3-BD: 0.68 ml/kg or 676 mg/kg as a 6.8 % aqueous
     solution (oral or i.v.). 3H-2B: 400 or 800 mg/kg as a 60% aqueous
     solution (i.v.)
* *
    Frequency of Treatment: Single Dose (oral or i.v.)
* *
    Duration of Test: 30 Hours
* *
     Sex: Male
* *
    Number/Dose Group: Not Identified
F020 255644
EOR
F002 27
F010 5.0
F004 1
F005 RS
F006 In the sBA-treatment phase of the study, blood concentrations of sBA and
     its metabolites MEK, 3H-2B, and 2,3-BD were measured. Blood sBA
     concentrations of 0.59 mg/ml peaked at 2 hours and declined to less than
     0.05 mg/ml at 16 hours. As th
F007 In the sBA-treatment phase of the study, blood concentrations of sBA and
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its metabolites MEK, 3H-2B, and 2,3-BD were measured. Blood sBA
     concentrations of 0.59 mg/ml peaked at 2 hours and declined to less than
     0.05 mg/ml at 16 hours. As the sBA concentration fell, the metabolite
     concentrations of MEK, 3H-2B, and 2,3-BD concentrations rose to maximums
     at 8, 12, and 18hr, respectively. The peak concentration of MEK was 0.78
    mg/ml, while that of 2,3-BD was 0.21 mg/ml. 3H-2B reached a peak
     concentration of 0.04 mg/ml. Total AUC values for sBA, MEK, 3H-2B, and
     2,3-BD were 3254 \pm 258, 9868 \pm 566, 443 \pm 93, and 3167 \pm 503 mg-hr/l,
    respectively.
    Following an oral dose of MEK, blood concentrations of MEK and its
    metabolites sBA, 3H-2B, and 2,3-BD were measured. Blood MEK
     concentrations of 0.95 mg/ml peaked at 4 hours and declined to less than
     0.07 mg/ml at 18 hours. As the MEK concentration fell, the end
    metabolite 2,3-BD rose to a maximum concentration of 0.26 mg/ml at 18
    hours. Peak concentrations of sBA and 3H-2B were 0.033 and 0.027 mg/ml,
     respectively. These were detected at 6 and 8 hr after the MEK
     administration. Total AUC values for MEK, sBA, 3H-2B, and 2,3-BD were
     10,899 \pm 824, 414 \pm 38, 382 \pm 38, and 3863 \pm 238 mg-hr/l, respectively.
F020 255638
EOR
F002 27
F010 5.0
F004 1
F005 TS
F006 2-Butanol (sec-Butanol; sBA) and metabolites 2-Butanone (Methyl ethyl
    ketone; MEK), 3-Hydroxy-2-Butanone (3H-2B), and 2,3-Butanediol (2,3-BD)
     (CAS Nos. 78-92-2 (sBA); 78-93-3 (MEK))
    Purity not identified.
F007 2-Butanol (sec-Butanol; sBA) and metabolites 2-Butanone (Methyl ethyl
    ketone; MEK), 3-Hydroxy-2-Butanone (3H-2B), and 2,3-Butanediol (2,3-BD)
     (CAS Nos. 78-92-2 (sBA); 78-93-3 (MEK))
     Purity not identified.
F020 255643
EOR
F002 27
F010 5.0
F004 2
F005 CL
F006 This study showed the initial metabolism of MEK follows both oxidative
     and reductive pathways to produce 3-hydroxy-2-butanone, 2,3-butanediol
     and 2-butanol.
F007 This study showed the initial metabolism of MEK follows both oxidative
     and reductive pathways to produce 3-hydroxy-2-butanone, 2,3-butanediol
     and 2-butanol.
F020 255648
EOR
F002 27
F010 5.0
F004 2
F005 RE
F006 DiVincenzo GD, Kaplan CJ and Dedinas J (1976). Characterization of the
     metabolites of Methyl n-Butyl Ketone, Methyl iso-Butyl Ketone, and Methyl
     Ethyl Ketone in guinea pig serum and their clearance. Toxicology and
     Applied Pharmacology 36
F007 DiVincenzo GD, Kaplan CJ and Dedinas J (1976). Characterization of the
     metabolites of Methyl n-Butyl Ketone, Methyl iso-Butyl Ketone, and Methyl
     Ethyl Ketone in guinea pig serum and their clearance. Toxicology and
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Applied Pharmacology 36, 511-522.
F020 255651
EOR
F002 27
F010 5.0
F004 2
F005 RT
F006 No circumstances occurred that would have affected the quality or
     integrity of the data. Test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
F007 No circumstances occurred that would have affected the quality or
     integrity of the data. Test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
F020 255650
EOR
F002 27
F010 5.0
F004 2
F005 RM
F006 Male guinea pigs ranging in weight from 250 - 450 grams were given a
     single i.p. dose of 450mg/kg MEK (study also investigated MnBK and MiBK -
     data not addressed here). Blood was collected by heart puncture from 4
     animals at each of the fol
F007 Male guinea pigs ranging in weight from 250 - 450 grams were given a
     single i.p. dose of 450mg/kg MEK (study also investigated MnBK and MiBK -
     data not addressed here). Blood was collected by heart puncture from 4
     animals at each of the following times after dose administration: 1, 2,
     4, 6, 8, 12, and 16 Hr. Only 1 sample was collected from each guinea pig.
     Serum was separated and refrigerated until assayed within 48 Hr. The
     concentrations of the ketones and their metabolites were measured in
     duplicate by direct on-column injection of undiluted serum into a Varian
     2100 Gas Chromatograph equipped with a flame ionization detector.
    Ketones and metabolites were quantitated from calibration curves prepared
     from pure standards. Greater than 90% of each ketone was distributed in
     the plasma fraction. Half-lives were estimated by extrapolating the
     linear portion of the decay curve to zero time.
F020 255647
EOR
F002 27
F010 5.0
F004 2
F005 RM
F006 Test Type: Metabolism and Pharmacokinetic Evaluation
     Frequency of Treatment: Single injection
* *
     Duration of Test: 16 Hours
    Number/Dose Group: 4 at each blood - sampling interval
F007 Test Type: Metabolism and Pharmacokinetic Evaluation
* *
     Frequency of Treatment: Single injection
* *
    Duration of Test: 16 Hours
* *
    Number/Dose Group: 4 at each blood - sampling interval
F020 255649
EOR
F002 27
F010 5.0
F004 2
F005 RS
F006 MEK: 2-Butanol, 3-hydroxy-2-butanone, and 2,3-butanediol were identified
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as metabolites in the serum of guinea pigs injected i.p. with MEK. The
    half-life of MEK in serum was 270 min. The clearance time of MEK was 12
     Hr. 2,3-Butanediol was c
F007 MEK: 2-Butanol, 3-hydroxy-2-butanone, and 2,3-butanediol were identified
     as metabolites in the serum of guinea pigs injected i.p. with MEK. The
    half-life of MEK in serum was 270 min. The clearance time of MEK was 12
    Hr. 2,3-Butanediol was cleared in 16 hours, as were the other two
    metabolites. The metabolism was described as oxidation via hydroxylation
     of the ? -1 carbon forming 3-hydroxy-2-butanone as the metabolite of MEK.
     Reduction occurred at the carbonyl group as expected forming the
     secondary alcohol, 2-butanol from MEK.
F020 255646
EOR
F002 27
F010 5.0
F004 2
F005 TS
F006 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)
     Purity: 98% with traces of 2-butanol
F007 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)
    Purity: 98% with traces of 2-butanol
F020 255645
EOR
F002 27
F010 5.0
F004 3
F005 CL
F006 Two elimination phases were detected for MEK in blood. The T% for the
     faster phase of elimination was 30 min and about 81 min for the slower
     phase. 2-3% of absorbed dose of MEK was eliminated by exhalation. Urinary
     excretion of unchanged ME
F007 Two elimination phases were detected for MEK in blood. The T% for the
     faster phase of elimination was 30 min and about 81 min for the slower
    phase. 2-3% of absorbed dose of MEK was eliminated by exhalation. Urinary
     excretion of unchanged MEK was 0.1% and excretion of the metabolite 2H3B
     was about 0.1%.
F020 255655
EOR
F002 27
F010 5.0
F004 3
F005 RE
F006 Liira J, Riihimaki V and Pfaffli P (1988). Kinetics of methyl ethyl
     ketone in man: absorption, distribution and elimination in inhalation
     exposure. Int.Arch. Occup. Environ. Health 60, 195-200.
F007 Liira J, Riihimaki V and Pfaffli P (1988). Kinetics of methyl ethyl
     ketone in man: absorption, distribution and elimination in inhalation
     exposure. Int.Arch. Occup. Environ. Health 60, 195-200.
F020 255658
EOR
F002 27
F010 5.0
F004 3
F005 RL
F006 No circumstances occurred that would have affected the quality or
     integrity of the data. Test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
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F007 No circumstances occurred that would have affected the quality or
     integrity of the data. Test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
F020 255657
EOR
F002 27
F010 5.0
F004 3
F005 RM
F006 Nine healthy male volunteers, 18-34 years of age (mean 23.7), weight
     65-81 kg (mean 74.3), height 172-190 cm (mean 181.8) and calculated
     surface area 1.81 - 2.09 m2 (mean 1.97) were exposed for 4 Hr to 200ppm
     MEK on 2 separate days at least
F007 Nine healthy male volunteers, 18-34 years of age (mean 23.7), weight
     65-81 kg (mean 74.3), height 172-190 cm (mean 181.8) and calculated
     surface area 1.81 - 2.09 m2 (mean 1.97) were exposed for 4 Hr to 200ppm
    MEK on 2 separate days at least 1 week apart. Venous blood samples were
     collected at 1-hr intervals during exposure and over 120 to 210 minutes
    post-exposure. In a supplementary study, follow-up elimination blood
     samples were collected in 2 persons until the next morning. One of the
     exposures constituted sedentary activity and the other encompassed three
     100 W ergometric exercise periods over minutes 5-15, 95-105 and 225-235
    during a total exposure of 240 minutes. Exhaled air samples were
     collected with a 2-way respirator mouthpiece into 4-liter polyester
     laminated aluminum-foil bags and analyzed immediately. Samples were
     collected at one-hour intervals during exposure and over 120-210 minutes
     thereafter. Urine samples were obtained at 2-hour intervals during the
     exposure day and in separate samples until the next morning. Whole
    venous blood was analyzed by gas chromatography, as were air samples.
    Peaks were compared to calibration curves prepared of known blood or air
     concentrations of MEK. 2-3BD in urine was analyzed according to a
     modification of the method of Robinson and Reive. The 2 peaks in the
     chromatograph (d,l-forms and meso-form) were summed for calculations.
F020 255654
EOR
F002 27
F010 5.0
F004 3
F005 RM
F006 Test Type: Metabolism and Pharmacokinetic Evaluation
     Frequency of Treatment: 2 Exposure periods at least 1 wk between
    Duration of Test: 4 Hr. Exposure
F007 Test Type: Metabolism and Pharmacokinetic Evaluation
     Frequency of Treatment: 2 Exposure periods at least 1 wk between
     Duration of Test: 4 Hr. Exposure
F020 255656
EOR
F002 27
F010 5.0
F004 3
F005 RS
F006 Pulmonary retention of MEK in the lungs remained constant throughout
     exposure with or without exercise. Relative uptake was 53% ± 2%.
     Estimated mean total pulmonary uptakes were 11.4 mmol (ventilation volume
     11 liters/min at rest) and 14.3
F007 Pulmonary retention of MEK in the lungs remained constant throughout
     exposure with or without exercise. Relative uptake was 53% ± 2%.
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Estimated mean total pulmonary uptakes were 11.4 mmol (ventilation volume
     11 liters/min at rest) and 14.3 mmol (ventilation volume 35 liters/min
     during the exercise). Blood MEK concentrations increased rapidly during
     the first hour of exposure (markedly faster when associated with
     exercise). Thereafter, concentrations increased slowly and linearly
     through 4 hr with sedentary activity and steeply during the exercise
    period at the end of the 4 hr exposure period. Two elimination phases
    were detected for MEK in blood. The calculated half-time for the faster
    phase of elimination (0-45 min post-exposure) was 30 min and about 81 min
     for the slower phase (60-320 min post-exposure). Owing to remarkable
     solubility of MEK, only 2-3% of absorbed dose was eliminated by
     exhalation. Urinary excretion of unchanged MEK was 0.1% and excretion of
     the metabolite 2H3B was about 0.1%. The authors speculated that the
     greater part of absorbed MEK is probably converted to products of
     intermediary metabolism, e.g., to acetate or acetoacetate via 3H2B.
F020 255653
EOR
F002 27
F010 5.0
F004 3
F005 TS
F006 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)
** Purity: Analytical Grade
F007 2-Butanone (Methyl ethyl ketone; MEK) (CAS No. 78-93-3)
    Purity: Analytical Grade
F020 255652
EOR
F002 27
F010 5.1.1
F004 1
F005 RE
F006 Environmental Health Criteria 65, World Health Organization,
F007 Environmental Health Criteria 65, World Health Organization,
     1987.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120141
EOR
F002 27
F010 5.1.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120142
EOR
F002 27
F010 5.1.1
F004 2
F005 SO
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F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120143
EOR
F002 27
F010 5.1.1
F004 3
F005 CL
F006 The acute oral LD50 for sBA in Fischer 344 rats is 2193 mg/kg.
F007 The acute oral LD50 for sBA in Fischer 344 rats is 2193 mg/kg.
F008 CLGETTS
F012 20
F020 120144
EOR
F002 27
F010 5.1.1
F004 3
F005 RE
F006 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell Research Report
     SBGR.86.108. Shell Rese
F007 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell Research Report
     SBGR.86.108. Shell Research Limited, Sittingbourne Research Centre.
F008 CLGETTS
F012 20
F020 120145
EOR
F002 27
F010 5.1.1
F004 3
F005 RL
F006 1 - Reliable study without restrictions. No circumstances occurred that
     would have affected the quality or integrity of the data.
F007 1 - Reliable study without restrictions. No circumstances occurred that
     would have affected the quality or integrity of the data.
F008 CLGETTS
F012 20
F020 120146
EOR
F002 27
F010 5.1.1
F004 3
F005 RM
F006 Fischer 344 rats were in the age range of 55-65 days when purchased and
     were 9 -11 weeks old when the study commenced. All animals were fasted
     for 18 hours prior to dosing and doses were altered by varying the volume
     dispensed from the syri
F007 Fischer 344 rats were in the age range of 55-65 days when purchased and
     were 9 -11 weeks old when the study commenced. All animals were fasted
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for 18 hours prior to dosing and doses were altered by varying the volume
     dispensed from the syringe. Body weights were recorded on Days 1, 7 and
     14. All surviving rats gained weight by the end of the 14-day observation
     period. The LD50's were calculated using probit analysis
F008 CLGETTS
F012 20
F020 120147
EOR
F002 27
F010 5.1.1
F004 3
F005 RM
F006 Route of Administration:
                              Oral Gavage
    Doses: 950 - 2400 mg/kg
     Volume Administered: Single dose volume varied by body weight
* *
     Post Dose Observation Period: Daily for 14 Days
* *
     Purity: 99.5%
F007 Route of Administration: Oral Gavage
    Doses: 950 - 2400 mg/kg
* *
    Volume Administered: Single dose volume varied by body weight
* *
    Post Dose Observation Period: Daily for 14 Days
* *
    Purity:
             99.5%
F008 CLGETTS
F012 20
F020 120148
EOR
F002 27
F010 5.1.1
F004 3
F005 RS
F006 2054 mg/kg (95% fiducial limits, 1283 - 4018 mg/kg) males,
     2328 mg/kg (95% fiducial limits, 1470 - 5428 mg/kg) females, and
     2193 mg/kg (95% fiducial limits, 1608 - 4146 mg/kg) combined.
F007 2054 mg/kg (95% fiducial limits, 1283 - 4018 mg/kg) males,
     2328 mg/kg (95% fiducial limits, 1470 - 5428 mg/kg) females, and
     2193 mg/kg (95% fiducial limits, 1608 - 4146 mg/kg) combined.
F008 CLGETTS
F012 20
F020 120149
EOR
F002 27
F010 5.1.1
F004 4
F005 CL
F006 The acute oral LD50 for sBA in Carworth-Wistar rats is 6.48 g/kg (5.73 -
F007 The acute oral LD50 for sBA in Carworth-Wistar rats is 6.48 g/kg (5.73 -
     7.32)
F008 CLGETTS
F012 20
F020 120150
EOR
F002 27
F010 5.1.1
F004 4
F005 RE
F006 Smyth, H. F. Jr., Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
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(1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F007 Smyth, H. F. Jr., Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
     (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F008 CLGETTS
F012 20
F020 120151
EOR
F002 27
F010 5.1.1
F004 4
F005 RL
F006 Smyth, H. F. Jr., Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
     (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F007 Smyth, H. F. Jr., Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
     (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F008 CLGETTS
F012 20
F020 120152
EOR
F002 27
F010 5.1.1
F004 4
F005 RM
F006 Route of Administration: Oral Gavage
     Doses: Logarithmic series
* *
     Dose Volume Administered: Single dose; 1-10 ml/rat
* *
     Post Dose Observation Period: 14 Days
F007 Route of Administration: Oral Gavage
* *
     Doses: Logarithmic series
* *
     Dose Volume Administered: Single dose; 1-10 ml/rat
* *
     Post Dose Observation Period: 14 Days
F008 CLGETTS
F012 20
F020 120153
EOR
F002 27
F010 5.1.1
F004 4
F005 RM
F006 The animals weighed 90 - 120 grams and were not fasted prior to dosing.
     The most probable LD50 value and the fiducial range were estimated by the
     method of Thompson using the tables of Weil.
F007 The animals weighed 90 - 120 grams and were not fasted prior to dosing.
     The most probable LD50 value and the fiducial range were estimated by the
     method of Thompson using the tables of Weil.
F008 CLGETTS
F012 20
F020 120154
EOR
F002 27
F010 5.1.1
F004 4
F005 RS
```

```
F006 6.48 g/kg (5.73 - 7.32)
F007 6.48 g/kg (5.73 - 7.32)
F008 CLGETTS
F012 20
F020 120155
EOR
F002 27
F010 5.1.1
F004 5
F005 CL
F006 The reported LD50 was 66 mmol/kg (4890 mg/kg).
F007 The reported LD50 was 66 mmol/kg (4890 \text{ mg/kg}).
F008 CLGETTS
F012 20
F020 120156
EOR
F002 27
F010 5.1.1
F004 5
F005 RE
F006 Munch, J.C. (1972). Aliphatic alcohols and alkyl esters: narcotic and
     lethal potencies to tadpoles and to rabbits. Ind. Med. 41:31-33.
F007 Munch, J.C. (1972). Aliphatic alcohols and alkyl esters: narcotic and
     lethal potencies to tadpoles and to rabbits. Ind. Med. 41:31-33.
F008 CLGETTS
F012 20
F020 120157
F:OR
F002 27
F010 5.1.1
F004 5
F005 RE
F006 Munch, J.C. and Schwartze, E.W. (1925). Narcotic and toxic potency of
     aliphatic alcohols upon rabbits. J Lab Clin Med, 10:985-996.
F007 Munch, J.C. and Schwartze, E.W. (1925). Narcotic and toxic potency of
     aliphatic alcohols upon rabbits. J Lab Clin Med, 10:985-996.
F008 CLGETTS
F012 20
F020 120158
EOR
F002 27
F010 5.1.1
F004 5
F005 RL
F006 3 - Not Reliable. Study is pre-GLP and documentation insufficient for
     assessment
F007 3 - Not Reliable. Study is pre-GLP and documentation insufficient for
     assessment
F008 CLGETTS
F012 20
F020 120159
EOR
F002 27
F010 5.1.1
F004 5
F005 RM
F006 Route of Administration: Oral Gavage
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Number of Animals/Dose: Unknown
    Doses: 14 or 66 mmol/kg; 1037 or 4890 mg/kg
* *
    Dose Volume Administered: < 50 ml
* *
     Post Dose Observation Period: 24 Hours
F007 Route of Administration: Oral Gavage
* *
    Number of Animals/Dose: Unknown
* *
    Doses: 14 or 66 mmol/kg; 1037 or 4890 mg/kg
    Dose Volume Administered: < 50 ml
    Post Dose Observation Period: 24 Hours
F008 CLGETTS
F012 20
F020 120160
EOR
F002 27
F010 5.1.1
F004 5
F005 RM
F006 Ten to 35 rabbits weighing between 1.5 and 2.5 kg were given "calculated"
     quantities of sBA at doses described as the "Narcotic Dose " or "Certain
     Lethal Dose" by oral gavage and observed for 24 hours.
F007 Ten to 35 rabbits weighing between 1.5 and 2.5 kg were given "calculated"
     quantities of sBA at doses described as the "Narcotic Dose " or "Certain
     Lethal Dose" by oral gavage and observed for 24 hours.
F008 CLGETTS
F012 20
F020 120161
EOR
F002 27
F010 5.1.1
F004 5
F005 RM
F006 The cited LD50 [66 mmol/kg, or 4890 mg/kg] is really the Certain Lethal
     Dose presented in the paper by Munch and Schwartze (1925). The Munch
     paper (1972) presents the rabbit data reported earlier in 1925 plus
     tadpole data. Because of the i
F007 The cited LD50 [66 mmol/kg, or 4890 mg/kg] is really the Certain Lethal
     Dose presented in the paper by Munch and Schwartze (1925). The Munch
    paper (1972) presents the rabbit data reported earlier in 1925 plus
     tadpole data. Because of the inconsistency in the definition of LD50 and
     Certain Lethal Dose, a CoR of 3 is assigned to both references.
F008 CLGETTS
F012 20
F020 120162
EOR
F002 27
F010 5.1.1
F004 5
F005 RS
F006 4890 mg/kg
F007 4890 mg/kg
F008 CLGETTS
F012 20
F020 120163
EOR
F002 27
F010 5.1.2
F004 1
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F005 CL
F006 Five of six exposed rats died within 14 days.
F007 Five of six exposed rats died within 14 days.
F008 CLGETTS
F012 20
F020 120164
EOR
F002 27
F010 5.1.2
F004 1
F005 RE
F006 Smyth, H. F. Jr., Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
     (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F007 Smyth, H. F. Jr., Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
     (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F008 CLGETTS
F012 20
F020 120165
EOR
F002 27
F010 5.1.2
F004 1
F005 RT
F006 2- Reliable with restrictions. Pre-GLP study documented, meets generally
     accepted scientific principles, acceptable for assessment.
F007 2- Reliable with restrictions. Pre-GLP study documented, meets generally
     accepted scientific principles, acceptable for assessment.
F008 CLGETTS
F012 20
F020 120166
EOR
F002 27
F010 5.1.2
F004 1
F005 RM
F006 Route of Administration: Inhalation
** Observation Period: Up to 14 days
F007 Route of Administration: Inhalation
    Observation Period: Up to 14 days
F008 CLGETTS
F012 20
F020 120167
EOR
F002 27
F010 5.1.2
F004 1
F005 RM
F006 The study consisted of exposing six male albino rats to a flowing stream
     of air approaching saturation with sBA vapors. The stream was prepared
    by proportioning pumps and nominal concentrations (not confirmed by
     analytical methods) were re
F007 The study consisted of exposing six male albino rats to a flowing stream
     of air approaching saturation with sBA vapors. The stream was prepared
    by proportioning pumps and nominal concentrations (not confirmed by
     analytical methods) were recorded.
                                        The study end point was the
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concentration yielding a fractional mortality among six rats within 14
     days.
F008 CLGETTS
F012 20
F020 120168
EOR
F002 27
F010 5.1.2
F004 1
F005 RS
F006 16,000 ppm sBA was lethal to 5 of 6 rats within 14 days
F007 16,000 ppm sBA was lethal to 5 of 6 rats within 14 days
F008 CLGETTS
F012 20
F020 120169
EOR
F002 27
F010 5.1.2
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120170
EOR
F002 27
F010 5.1.2
F004 2
F005 CL
F006 One of six exposed rats died within 14 days.
F007 One of six exposed rats died within 14 days.
F008 CLGETTS
F012 20
F020 120171
EOR
F002 27
F010 5.1.2
F004 2
F005 RE
F006 Mellon (Mellon Institute of Industrial Research). (1951). "Project Report
     Number 14-78". Union Carbide Corporation, Danbury, CT, USA.
F007 Mellon (Mellon Institute of Industrial Research). (1951). "Project Report
     Number 14-78". Union Carbide Corporation, Danbury, CT, USA.
F008 CLGETTS
F012 20
F020 120172
EOR
F002 27
F010 5.1.2
F004 2
F005 RL
F006 2- Reliable with restrictions. Pre-GLP study documented, meets generally
     accepted scientific principles, acceptable for assessment.
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F007 2- Reliable with restrictions. Pre-GLP study documented, meets generally
     accepted scientific principles, acceptable for assessment.
F008 CLGETTS
F012 20
F020 120173
EOR
F002 27
F010 5.1.2
F004 2
F005 RM
F006 Route of Administration: Inhalation
    Observation Period: Up to 14 days
F007 Route of Administration: Inhalation
     Observation Period: Up to 14 days
F008 CLGETTS
F012 20
F020 120174
EOR
F002 27
F010 5.1.2
F004 2
F005 RM
F006 The study consisted of exposing six male albino rats to a flowing stream
     of air with sBA vapors. The stream was prepared by proportioning pumps
     and nominal concentrations (not confirmed by analytical methods) were
     recorded.
                 The study end
F007 The study consisted of exposing six male albino rats to a flowing stream
     of air with sBA vapors. The stream was prepared by proportioning pumps
     and nominal concentrations (not confirmed by analytical methods) were
                 The study end point was the concentration yielding a
     fractional mortality among six rats within 14 days.
F008 CLGETTS
F012 20
F020 120175
EOR
F002 27
F010 5.1.2
F004 2
F005 RS
F006 8,000 ppm sBA was lethal to 1 of 6 rats within 14 days
F007 8,000 ppm sBA was lethal to 1 of 6 rats within 14 days
F008 CLGETTS
F012 20
F020 120176
EOR
F002 27
F010 5.1.2
F004 3
F005 CL
F006 Five of five exposed female rats died following a single 7 hour exposure.
F007 Five of five exposed female rats died following a single 7 hour exposure.
F008 CLGETTS
F012 20
F020 120177
EOR
F002 27
F010 5.1.2
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F004 3
F005 RE
F006 Nelson, B. K., Brightwell, W. S., Khan, A., Burg, J. R., and Goad, P. T.
     (1989). Lack of selective developmental toxicity of three butanol
     isomers administered by inhalation to rats. Fundam Appl Toxicol,
     12:469-479.
F007 Nelson, B. K., Brightwell, W. S., Khan, A., Burg, J. R., and Goad, P. T.
     (1989). Lack of selective developmental toxicity of three butanol
     isomers administered by inhalation to rats. Fundam Appl Toxicol,
     12:469-479.
F008 CLGETTS
F012 20
F020 120178
EOR
F002 27
F010 5.1.2
F004 3
F005 RL
F006 2 - Reliable. Pilot study with acceptable restrictions.
F007 2 - Reliable. Pilot study with acceptable restrictions.
F008 CLGETTS
F012 20
F020 120179
EOR
F002 27
F010 5.1.2
F004 3
F005 RM
F006 Study Type: Acute Inhalation (Pilot for Teratology study)
     Purity: >or= 99 %
     Route of Administration: Inhalation
     Observation Period: Unknown
F007 Study Type: Acute Inhalation (Pilot for Teratology study)
* *
     Purity: >or= 99 %
     Route of Administration: Inhalation
     Observation Period: Unknown
F008 CLGETTS
F012 20
F020 120180
EOR
F002 27
F010 5.1.2
F004 3
F005 RM
F006 The study consisted of exposing five female rats to a flowing stream of
     air with sBA vapors. The stream was prepared by proportioning pumps. The
     study end point was the concentration yielding a fractional mortality.
F007 The study consisted of exposing five female rats to a flowing stream of
     air with sBA vapors. The stream was prepared by proportioning pumps. The
     study end point was the concentration yielding a fractional mortality.
F008 CLGETTS
F012 20
F020 120181
EOR
F002 27
F010 5.1.2
F004 3
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F005 RS
F006 10,000 ppm sBA was lethal to 5 of 5 rats
F007 10,000 ppm sBA was lethal to 5 of 5 rats
F008 CLGETTS
F012 20
F020 120182
EOR
F002 27
F010 5.1.2
F004 4
F005 CL
F006 Doses from 3,300 to 19,800 ppm induced narcosis but not death.
F007 Doses from 3,300 to 19,800 ppm induced narcosis but not death.
F008 CLGETTS
F012 20
F020 120183
EOR
F002 27
F010 5.1.2
F004 4
F005 RE
F006 Starrek, E. (1938). Dissertation, Wirzburg.
F007 Starrek, E. (1938). Dissertation, Wirzburg.
F008 CLGETTS
F012 20
F020 120184
EOR
F002 27
F010 5.1.2
F004 4
F005 RL
F006 4 - Documentation insufficient for assessment. Non-guideline pre-GLP
F007 4 - Documentation insufficient for assessment. Non-guideline pre-GLP
     study.
F008 CLGETTS
F012 20
F020 120185
EOR
F002 27
F010 5.1.2
F004 4
F005 RM
F006 1650 ppm: no signs of intoxication after 420 minutes.
     3300 ppm: ataxia in 51-100 minutes, prostration in 120-180 minutes,
     narcosis in 3000 minutes, no deaths
* *
     19,800 ppm: ataxia in 7-8 minutes, prostration in 12-20 minutes, narcosis
     in 40 m
F007 1650 ppm: no signs of intoxication after 420 minutes.
     3300 ppm: ataxia in 51-100 minutes, prostration in 120-180 minutes,
     narcosis in 3000 minutes, no deaths
     19,800 ppm: ataxia in 7-8 minutes, prostration in 12-20 minutes, narcosis
     in 40 minutes, no deaths
F008 CLGETTS
F012 20
F020 120186
EOR
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F002 27
F010 5.1.2
F004 4
F005 RM
F006 Route of Administration: Inhalation
     Doses/time: 1650, 3300, 19800 ppm/decreasing lengths of time
     Post Dose Observation Period: Various
F007 Route of Administration: Inhalation
    Doses/time: 1650, 3300, 19800 ppm/decreasing lengths of time
     Post Dose Observation Period: Various
F008 CLGETTS
F012 20
F020 120187
EOR
F002 27
F010 5.1.2
F004 4
F005 RS
F006 Not Defined
F007 Not Defined
F008 CLGETTS
F012 20
F020 120188
EOR
F002 27
F010 5.1.3
F004 1
F005 CL
F006 The acute percutaneous LD50 of undiluted sBA in rats was > 2000 mg/kg.
F007 The acute percutaneous LD50 of undiluted sBA in rats was > 2000 mg/kg.
F008 CLGETTS
F012 20
F020 120189
EOR
F002 27
F010 5.1.3
F004 1
F005 RE
F006 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Rese
F007 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Research Limited, Sittingbourne Research Centre.
F008 CLGETTS
F012 20
F020 120190
EOR
F002 27
F010 5.1.3
F004 1
F005 RL
F006 1 - Reliable without Restrictions. No circumstances occurred that would
     have affected the quality or integrity of the data.
F007 1 - Reliable without Restrictions. No circumstances occurred that would
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have affected the quality or integrity of the data.
F008 CLGETTS
F012 20
F020 120191
EOR
F002 27
F010 5.1.3
F004 1
F005 RM
F006 Fischer 344 rats were in the age range of 55-65 days when purchased and
     were 9 -11 weeks old when the study commenced. The day before dosing the
     animals, approximately 60% of the dorsal hair was closely shorn with fine
     electric clippers, a
F007 Fischer 344 rats were in the age range of 55-65 days when purchased and
     were 9 -11 weeks old when the study commenced. The day before dosing the
     animals, approximately 60% of the dorsal hair was closely shorn with fine
     electric clippers, and the rats and the rats were weighed on the day of
    dosing. Varying the volume dispensed from the syringe altered doses.
    Application was made at room temperature and the test material was
     covered with aluminum foil and held in place by a double over-wrap of
    waterproof adhesive tape. The rats were individually housed for the next
     24 hours. Food was withheld, but water was available ad libitum. At the
    end of the 24-hour period, the tape and foil were removed and the skin
    was washed with warm dilute detergent solution and then dried.
    animals were observed for signs of toxicity for 14 days after dosing.
    Body weights were recorded on Days 1, 7 and 14. None of the rats died.
    There were no overt signs of toxicity and all rats gained weight relative
     to their day 1body weight.
F008 CLGETTS
F012 20
F020 120192
EOR
F002 27
F010 5.1.3
F004 1
F005 RM
F006 Route of Administration: Dermal
    Doses/time: 2000 mg/kg single application / 24-hour occlusive patch
* *
     Post Dose Observation Period: Daily for 14 Days
* *
     Purity: 99.5%
F007 Route of Administration: Dermal
    Doses/time: 2000 mg/kg single application / 24-hour occlusive patch
     Post Dose Observation Period: Daily for 14 Days
* *
    Purity: 99.5%
F008 CLGETTS
F012 20
F020 120193
EOR
F002 27
F010 5.1.3
F004 1
F005 RS
F006 > 2000 \text{ mg/kg}
F007 > 2000 \, mg/kg
F008 CLGETTS
F012 20
F020 120194
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EOR
F002 27
F010 5.1.4
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120195
EOR
F002 27
F010 5.10
F004 1
F005 RE
F006 Environmental Health Criteria 65. Butanols: Four Isomers.
     World Health Organization, 1987.
F007 Environmental Health Criteria 65. Butanols: Four Isomers.
    World Health Organization, 1987.
F008 HEDSET
F009 22-10-1992
F012 20
F020 120203
EOR
F002 27
F010 5.10
F004 1
F005 RM
F006 Excessive exposure by inhalation may result in headache,
     dizziness, drowsiness and narcosis. No adverse systemic
* *
     effects due to exposure to 2-butanol have been reported in
* *
F007 Excessive exposure by inhalation may result in headache,
     dizziness, drowsiness and narcosis. No adverse systemic
     effects due to exposure to 2-butanol have been reported in
    man.
F008 HEDSET
F009 22-10-1992
F012 20
F020 120204
EOR
F002 27
F010 5.10
F004 1
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120205
EOR
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F002 27
F010 5.11
F004 1
F005 CL
F006 sBA caused a concentration-dependent decrease in respiratory tidal volume
     in normal and cannulated mice and acts as a respiratory irritant.
F007 sBA caused a concentration-dependent decrease in respiratory tidal volume
     in normal and cannulated mice and acts as a respiratory irritant.
F008 CLGETTS
F012 20
F020 120196
EOR
F002 27
F010 5.11
F004 1
F005 RE
F006 Hansen, L.F., and Nielsen, G.D. (1994). Sensory irritation, pulmonary
     irritation and structure-activity relationships of alcohols. Toxicol.,
     88:81-99.
F007 Hansen, L.F., and Nielsen, G.D. (1994). Sensory irritation, pulmonary
     irritation and structure-activity relationships of alcohols. Toxicol.,
     88:81-99.
F008 CLGETTS
F012 20
F020 120197
EOR
F002 27
F010 5.11
F004 1
F005 RL
F006 1 - Reliable without Restrictions. No circumstances occurred that would
    have affected the quality or integrity of the data.
F007 1 - Reliable without Restrictions. No circumstances occurred that would
    have affected the quality or integrity of the data.
F008 CLGETTS
F012 20
F020 120198
EOR
F002 27
F010 5.11
F004 1
F005 RM
F006 Respiratory rates of CF-1 mice (mean weight 25 g.) were determined prior
     to exposure. SBA was evaporated, diluted with room air and fed into a 3.3
     liter chamber at an airflow rate of 18.3 to 26.9 l/min. Each animal was
     placed in a body ple
F007 Respiratory rates of CF-1 mice (mean weight 25 g.) were determined prior
     to exposure. SBA was evaporated, diluted with room air and fed into a 3.3
     liter chamber at an airflow rate of 18.3 to 26.9 l/min. Each animal was
    placed in a body plethysmograph attached to the exposure chamber so that
     the head of the animal protruded into the chamber. The respiratory rate
    and the relative tidal volume were obtained by attaching a pressure
     transducer to each plethysmograph. The results were expressed as
    percentage of pre-exposure values, obtained from a 10-minute control
    period. The exposure was 30 minutes, followed by a 20-minute recovery
    period during which the respiratory pattern and respiratory rate were
    monitored continuously. Respiration rates and tidal volumes were compared
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between normal and cannulated mice to determine the on-set and extent of
     respiratory irritation. In the cannulated mice, the onset of the decrease
     in respiration was slower than that of the normal mice indicating that
     sBA was acting as a respiratory irritant.
F008 CLGETTS
F012 20
F020 120199
EOR
F002 27
F010 5.11
F004 1
F005 RM
F006 Species/Strain: Mouse / CF-1
     Sex: Male
     Number/Sex/Dose: 10
* *
     Vehicle: None
* *
     Route of Administration: Inhalation
* *
     Doses: 2800, 5600, 10100 or 15300 ppm normal mice;
* *
     10500 or 14000 ppm cannulated mice
* *
     Time of Exposure: 30 minute
* *
     Post Dose
F007 Species/Strain: Mouse / CF-1
* *
     Sex: Male
* *
     Number/Sex/Dose: 10
* *
     Vehicle: None
* *
     Route of Administration: Inhalation
* *
    Doses: 2800, 5600, 10100 or 15300 ppm normal mice;
* *
     10500 or 14000 ppm cannulated mice
     Time of Exposure: 30 minute
* *
     Post Dose Observation Period: 20 minute
* *
    Method: Alarie Test
* *
     GLP: Unknown
* *
     Year: 1994
F008 CLGETTS
F012 20
F020 120200
EOR
F002 27
F010 5.11
F004 1
F005 RS
F006 640 ppm RD0 Threshold Concentration at 2 minutes;
     11800 ppm RD50 at 10 minutes
F007 640 ppm RD0 Threshold Concentration at 2 minutes;
     11800 ppm RD50 at 10 minutes
F008 CLGETTS
F012 20
F020 120201
EOR
F002 27
F010 5.11
F004 1
F005 TS
F006 sec-Butanol (sBA)
F007 sec-Butanol (sBA)
F008 CLGETTS
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F012 20

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F020 120202
EOR
F002 27
F010 5.2.1
F004 1
F005 RE
F006 Environmental Health Criteria 65, Butanols: Four Isomers.
     World Health Organization, 1987.
F007 Environmental Health Criteria 65, Butanols: Four Isomers.
    World Health Organization, 1987.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120206
EOR
F002 27
F010 5.2.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120207
F:OR
F002 27
F010 5.2.1
F004 2
F005 CL
F006 sBA caused no skin reaction and is not a skin irritant in rabbits.
F007 sBA caused no skin reaction and is not a skin irritant in rabbits.
F008 CLGETTS
F012 20
F020 120208
EOR
F002 27
F010 5.2.1
F004 2
F005 RE
F006 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Rese
F007 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Research Limited, Sittingbourne Research Centre.
F008 CLGETTS
F012 20
F020 120209
EOR
F002 27
F010 5.2.1
F004 2
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F005 RT
F006 1 - Reliable without Restrictions. No circumstances occurred that would
     have affected the quality or integrity of the data
F007 1 - Reliable without Restrictions. No circumstances occurred that would
    have affected the quality or integrity of the data
F008 CLGETTS
F012 20
F020 120210
EOR
F002 27
F010 5.2.1
F004 2
F005 RM
F006 New Zealand White rabbits, 3 - 6 months of age, were acclimated to the
     laboratory conditions for at least 2 weeks prior to experiment
     initiation. The hair of the dorsal back between the shoulders and
     hindquarters of three male and 3 female
F007 New Zealand White rabbits, 3 - 6 months of age, were acclimated to the
     laboratory conditions for at least 2 weeks prior to experiment
     initiation. The hair of the dorsal back between the shoulders and
    hindquarters of three male and 3 female rabbits, 4 - 9 months of age at
    dosing, was closely shorn with fine electric clippers. A test site was
     selected and a 2 cm x 2 cm lint patch with 0.5 ml of sBA was applied to
     it. The patch and surrounding skin were covered by a single layer of
    gauze and held in place with an elastic adhesive bandage. The wrapping
    and patch were removed after 4 hours and the skin was not washed. The
     site was examined and scored for erythema and edema on a graded scale 0 -
     4. Observations were made 30 minutes after the removal of the patch and
    at 24, 48, and 72 hours and 7 days following dosing. The mean scores for
     each rabbit at each observation time were calculated. There were no skin
     reactions following the application of sBA to rabbit skin for 4 hours.
     The test material is therefore not a skin irritant in rabbits.
F008 CLGETTS
F012 20
F020 120211
EOR
F002 27
F010 5.2.1
F004 2
F005 RM
F006 Purity: 99.5%
    Sex: Male and Female
* *
    Vehicle: None
* *
    Route of Administration: Dermal
* *
    Doses: 0.5 ml
* *
    Doses/Time: Single application
* *
     Post Dose Observation Period: After removal of patch: 0.5 hours; 24, 48,
     72 hours and Day 7 after dosing.
F007 Purity: 99.5%
* *
     Sex: Male and Female
* *
    Vehicle: None
* *
    Route of Administration: Dermal
* *
    Doses: 0.5 ml
* *
    Doses/Time: Single application
     Post Dose Observation Period: After removal of patch: 0.5 hours; 24, 48,
     72 hours and Day 7 after dosing.
F008 CLGETTS
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F012 20
F020 120212
EOR
F002 27
F010 5.2.1
F004 2
F005 RS
F006 PII = 0
F007 PII = 0
F008 CLGETTS
F012 20
F020 120213
EOR
F002 27
F010 5.2.2
F004 1
F005 CL
F006 sBA caused moderate damage and/or corneal opacity in rabbits.
F007 sBA caused moderate damage and/or corneal opacity in rabbits.
F008 CLGETTS
F012 20
F020 120214
EOR
F002 27
F010 5.2.2
F004 1
F005 RE
F006 Carpenter, C.P., and Smith, H. F. Jr,. (1946). Chemical Burns of the
     Rabbit Cornea. Am. J. Ophth. 29:1363.
F007 Carpenter, C.P., and Smith, H. F. Jr,. (1946). Chemical Burns of the
     Rabbit Cornea. Am. J. Ophth. 29:1363.
F008 CLGETTS
F012 20
F020 120215
EOR
F002 27
F010 5.2.2
F004 1
F005 RE
F006 Smyth, H. F. Jr, Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
     (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F007 Smyth, H. F. Jr, Carpenter, C. P., Weil, C. S., and Pozzani, U. C.
     (1954). Range-finding toxicity data: List V. Arch Ind Hyg Occup Med,
     10:61-68.
F008 CLGETTS
F012 20
F020 120216
EOR
F002 27
F010 5.2.2
F004 1
F005 RL
F006 2 - Reliable with restrictions: data are from a collection of data.
F007 2 - Reliable with restrictions: data are from a collection of data.
F008 CLGETTS
F012 20
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F020 120217
EOR
F002 27
F010 5.2.2
F004 1
F005 RM
F006 Eye injury in rabbits records the degree of corneal narcosis from various
     volumes and concentrations of chemical, as detailed in Carpenter and
     Smyth (1946). Grade 1 indicates at most a very small area of narcosis
     resulting from 0.5 ml of u
F007 Eye injury in rabbits records the degree of corneal narcosis from various
     volumes and concentrations of chemical, as detailed in Carpenter and
     Smyth (1946). Grade 1 indicates at most a very small area of narcosis
     resulting from 0.5 ml of undiluted chemical in the eye; Grade 5 indicates
     a so-called severe burn from 0.005 ml, and Grade 10 indicates a severe
     burn from 0.5 ml of a 1% solution in water or propylene glycol.
F008 CLGETTS
F012 20
F020 120218
EOR
F002 27
F010 5.2.2
F004 1
F005 RM
F006 Sex: Male
     Vehicle: None
* *
     Route of Administration: Ocular application into the conjunctival sac of
     one eye
     Control: Untreated eye
* *
     Dose: Single application of undiluted sBA
* *
     Volume: 0.02 and 0.1 ml
* *
     Post Dose Observation Period: 24 Hours
F007 Sex: Male
* *
     Vehicle: None
    Route of Administration: Ocular application into the conjunctival sac of
     one eye
* *
    Control: Untreated eye
* *
    Dose: Single application of undiluted sBA
    Volume: 0.02 and 0.1 ml
* *
     Post Dose Observation Period: 24 Hours
F008 CLGETTS
F012 20
F020 120219
EOR
F002 27
F010 5.2.2
F004 1
F005 RS
F006 Irritating (score 4/10); Severe injury from 0.1 ml, minor from 0.02 ml;
     Grade 4 eye injury.
F007 Irritating (score 4/10); Severe injury from 0.1 ml, minor from 0.02 ml;
     Grade 4 eye injury.
F008 CLGETTS
F012 20
F020 120220
EOR
F002 27
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F010 5.2.2
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120221
EOR
F002 27
F010 5.2.2
F004 2
F005 CL
F006 sBA caused moderate conjunctival inflammation in all six rabbits with
     slight transitory iritic damage and/or corneal opacity in three rabbits.
     One rabbit developed intense, extensive corneal opacity and complete loss
     of iritic response.
F007 sBA caused moderate conjunctival inflammation in all six rabbits with
     slight transitory iritic damage and/or corneal opacity in three rabbits.
     One rabbit developed intense, extensive corneal opacity and complete loss
     of iritic response. The test material was there for corrosive to the
    rabbit eye. On administration of sBA, there was a moderate initial pain
    response.
F008 CLGETTS
F012 20
F020 120222
EOR
F002 27
F010 5.2.2
F004 2
F005 RE
F006 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Rese
F007 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Research Limited, Sittingbourne Research Centre.
F008 CLGETTS
F012 20
F020 120223
EOR
F002 27
F010 5.2.2
F004 2
F005 RL
F006 1 - Reliable without Restrictions. No circumstances occurred that would
    have affected the quality or integrity of the data.
F007 1 - Reliable without Restrictions. No circumstances occurred that would
    have affected the quality or integrity of the data.
F008 CLGETTS
F012 20
F020 120224
```

```
EOR
F002 27
F010 5.2.2
F004 2
F005 RM
F006 New Zealand White rabbits, 3 - 6 months of age, were acclimated to the
     laboratory conditions for at least 2 weeks prior to experiment
     initiation. Three male and 3 female rabbits were 4 - 9 months of age at
     dosing. The day before testing, t
F007 New Zealand White rabbits, 3 - 6 months of age, were acclimated to the
     laboratory conditions for at least 2 weeks prior to experiment
     initiation. Three male and 3 female rabbits were 4 - 9 months of age at
     dosing. The day before testing, the eyes were carefully examined for any
     damage and any animals showing damage were replaced. A single dose of
     0.1 ml of sBA was placed into the lower conjunctival sac of one eye and
     the lids held together for a few seconds to prevent loss of material.
     The eyes were not washed. The reactions of the animal were observed and
     the initial pain responses graded on a 1 - 6 scale, no pain to very sever
     initial pain. Visual assessments were made 1, 5-7, 24, 28, and 72 hours
     and 7 Days postdosing. Irritancy was scored for the cornea, iris, and
    conjunctiva using standard scores. Any corneal damage visualization was
    aided by instillation of one drop of 2% fluorescein solution. The degree
     of irritation was classified using a scheme based on the OECD Data
     Interpretation Guide (1984). The instillation of undiluted sBA resulted
     in moderate initial pain. The conjunctival redness, chemosis and
    discharge, corneal opacity and damage to the iris were assessed and mean
     scores calculated. All rabbits had moderate conjunctival inflammation
    with some discharge within 1 hour of dosing. The swelling and discharge
     largely cleared by four hours but the redness persisted in 3 rabbits for
     7 days. These 3 animals, and another, had impaired iritic response
     and/or slight corneal opacity between 24 and 72 hours postdosing. The
     effect had cleared by 7 days in 3 rabbits. The severely affected rabbit
    had a completely opaque cornea and no iritic response. It was humanely
     terminated since recovery was deemed not possible. The remaining rabbits
     were retained and by day 14 all ocular effects had cleared. In view of
     the responses, sBA is classified as corrosive to rabbit eyes (OECD, 1984).
F008 CLGETTS
F012 20
F020 120225
EOR
F002 27
F010 5.2.2
F004 2
F005 RM
F006 Study Type: Ocular Irritation (Draize type)
     Purity: 99.5%
* *
     Sex: Male and Female
* *
    Vehicle: None
* *
    Route of Administration: Ocular application into the conjunctival sac of
     one eye
* *
     Control: Untreated Eye
     Dose: Single application of neat ma
F007 Study Type: Ocular Irritation (Draize type)
* *
     Purity: 99.5%
* *
     Sex: Male and Female
* *
    Vehicle: None
    Route of Administration: Ocular application into the conjunctival sac of
```

```
Control: Untreated Eye
* *
     Dose: Single application of neat material
* *
    Volume: 0.1 ml
    Post Dose Observation Period: 1, 5-7, 24, 28, and 72 hours and 7 Days
F008 CLGETTS
F012 20
F020 120226
EOR
F002 27
F010 5.2.2
F004 2
F005 RS
F006 Corrosive
F007 Corrosive
F008 CLGETTS
F012 20
F020 120227
EOR
F002 27
F010 5.3
F004 1
F005 CL
F006 In the guinea pig maximization test of Magnusson and Kligman, sBA did not
     induce positive responses in any of the 20 test animals at 24 or 48 hours
     after removal of the challenge patches. sBA is therefore not a dermal
     sensitizer in guinea
F007 In the guinea pig maximization test of Magnusson and Kligman, sBA did not
     induce positive responses in any of the 20 test animals at 24 or 48 hours
     after removal of the challenge patches. sBA is therefore not a dermal
     sensitizer in guinea pigs.
F008 CLGETTS
F012 20
F020 120228
EOR
F002 27
F010 5.3
F004 1
F005 RE
F006 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Rese
F007 Price, J. B. (1986). Toxicology of industrial chemicals (solvents): The
     acute oral and percutaneous toxicity, skin and eye irritancy and skin
     sensitizing potential of secondary butyl alcohol. Shell research Report
     SBGR.86.108. Shell Research Limited, Sittingbourne Research Centre.
F008 CLGETTS
F012 20
F020 120229
EOR
F002 27
F010 5.3
F004 1
F005 RL
F006 1 - Reliable without Restrictions. No circumstances occurred that would
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one eye

have affected the quality or integrity of the data. F007 1 - Reliable without Restrictions. No circumstances occurred that would have affected the quality or integrity of the data. F008 CLGETTS F012 20 F020 120230 EOR F002 27 F010 5.3 F004 1 F005 RM F006 Guinea pigs in the weight range of 300-370 grams were purchased from a commercial supplier and acclimated to the laboratory for at least 2 weeks. The skin sensitization potential of sBA was assessed using the guinea pig maximization test o F007 Guinea pigs in the weight range of 300-370 grams were purchased from a commercial supplier and acclimated to the laboratory for at least 2 weeks. The skin sensitization potential of sBA was assessed using the guinea pig maximization test of Magnusson and Kligman. The test was accomplished in 2 stages: rangefinding and definitive test. * * Rangefinding: * * The purpose of the rangefinding studies was to determine the concentrations of test material to be used for intradermal induction, topical induction and topical challenge. Two males and two female guinea pigs were closely shorn in the shoulder region using electric clippers followed by an electric razor and 0.1 ml of several dilutions (0.05, 0.1, 0.5 and 1.0% [m/v] in corn oil) of sBA injected intradermally each side of the midline. The animals were observed over the next few days to determine the maximum concentration that could be tolerated without causing untoward toxicity. Three further groups of two males and two females had their flanks closely shorn and 0.3 ml of several dilutions (25%, 50%, and 75% [m/v] in corn oil) of sBA were applied to a 4 cm x 4 cm Whatman Number 3 filter paper patches. The patches were placed on the flanks and held in place with a "Sleek" adhesive tape patch, then covered with a "Poroplast" elastic adhesive bandage for 24 hours. The bandages were removed and the animals were examined for signs of irritation and scored using a 4-point scale (0, 1, 2, and 3). The concentration used for topical induction was the one that scored 1 for irritation and the concentration for challenge was the one that was 0, a non-irritant. * * Definitive: This test was conducted using a group of ten male and ten female guinea pigs together with a control group of five males and five females. test consisted of two stages: a) induction by intradermal injection and topical application, and, 2) topical challenge. The following concentrations were selected for the definitive study: Intradermal induction, 0.1% (m/v) in corn oil; Topical induction (one week following intradermal challenge), 50% (m/v) in corn oil and 25% (m/v) in corn oil. Topical challenge was carried out two weeks following the topical induction by applying 0.1 ml of the diluted sBA to a semi-occluded challenge patch and removing it 24 hours later. The erythema resulting from the topical challenge was scored on a 4-point scale (0, 1, 2, and 3) immediately on removal of the challenge patches and 24 and 48 hours latter. None of the twenty test animals showed any positive response at either 24 or 48 hours after the removal of the challenge patches. It was

concluded that sBA was not a dermal sensitizer in guinea pigs.

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F008 CLGETTS
F012 20
F020 120231
EOR
F002 27
F010 5.3
F004 1
F005 RM
F006 Purity: 99.5%
* *
     Sex: Male and Female
* *
     Route of Administration: Intradermal and topical induction; topical
     challenge
* *
     Doses:
* *
     Intradermal Induction:
* *
     Test Doses:
* *
     2 injections (0.1 ml) of Freund's Complete Adjuvant (FCA)
* *
     2 injections (0.1 ml
F007 Purity: 99.5%
* *
* *
     Sex: Male and Female
* *
* *
     Route of Administration: Intradermal and topical induction; topical
     challenge
* *
* *
     Doses:
     Intradermal Induction:
* *
     Test Doses:
* *
     2 injections (0.1 ml) of Freund's Complete Adjuvant (FCA)
* *
     2 injections (0.1 ml) of sBA in corn oil
* *
     2 injections (0.1 ml) of sBA in 50:50 FCA:corn oil
* *
     Control Doses:
* *
     2 injections (0.1 ml) of Freund's Complete Adjuvant (FCA)
                                                              2 injections (0.1
     ml) of corn oil
* *
     2 injections (0.1 ml) of 50:50 FCA:corn oil
     Topical Induction:
* *
     Application of 0.3 ml 50% sBA (m/v) in corn oil and, 25% sBA (m/v) in
     corn oil covered for 48 hours.
* *
     Topical Challenge:
* *
     Application 0.1 ml of diluted sBA to a semi-occluded challenge patch and
     removing it 24 hours later.
* *
     Post Dose Observation Period: Immediately, 24 and 48 hours following
     challenge
F008 CLGETTS
F012 20
F020 120232
EOR
F002 27
F010 5.3
F004 1
F006 sBA is not a dermal sensitizer in guinea pigs.
F007 sBA is not a dermal sensitizer in guinea pigs.
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```
F008 CLGETTS
F012 20
F020 120233
EOR
F002 27
F010 5.3
F004 2
F005 CL
F006 In the guinea pig maximization test of Magnusson and Kligman, sBA is not
     a dermal sensitizer.
F007 In the guinea pig maximization test of Magnusson and Kligman, sBA is not
     a dermal sensitizer.
F008 CLGETTS
F012 20
F020 120234
EOR
F002 27
F010 5.3
F004 2
F005 RE
F006 Elf Atochem, Centre International de Toxicologie. 1997. Skin
     sensitization in guinea pigs (Maximization method of Magnusson, B. and
     Kligman, A.M.). Report # 14861 TSG.
F007 Elf Atochem, Centre International de Toxicologie. 1997.
     sensitization in guinea pigs (Maximization method of Magnusson, B. and
     Kligman, A.M.). Report # 14861 TSG.
F008 CLGETTS
F012 20
F020 120235
EOR
F002 27
F010 5.3
F004 2
F005 RL
F006 1 - Reliable without Restrictions.
F007 1 - Reliable without Restrictions.
F008 CLGETTS
F012 20
F020 120236
EOR
F002 27
F010 5.3
F004 2
F005 RM
F006 Age: ~ 3 months
* *
* *
     Sex: Male and Female
* *
* *
     Weight (initial): Males 335 \pm 17 g; Females 332 \pm 19 g
* *
* *
     Doses:
* *
     Intradermal Induction:
* *
     Test Doses:
* *
     Injections: sBA in paraffin oil (5% w/w)
* *
     sBA in FCA (Freund's Complete Adjuvant)
* *
     Control Doses:
* *
     In
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F007 Age: ~ 3 months
* *
* *
     Sex: Male and Female
* *
* *
     Weight (initial): Males 335 \pm 17 q; Females 332 \pm 19 q
* *
* *
* *
     Intradermal Induction:
* *
     Test Doses:
* *
     Injections: sBA in paraffin oil (5% w/w)
* *
     sBA in FCA (Freund's Complete Adjuvant)
     Control Doses:
* *
     Injections: Paraffin oil
* *
     Freund's Complete Adjuvant (FCA)
* *
     Topical Induction:
* *
     Application: sBA (undiluted) covered for 48 hours
* *
     Topical Challenge:
* *
     Application: sBA (undiluted) covered for 24 hours
* *
* *
     Post Dose Observation Period: Skin reactions were evaluated at
     approximately 24 and 48 hours.
F008 CLGETTS
F012 20
F020 120237
EOR
F002 27
F010 5.3
F004 2
F005 RM
F006 Clinical Signs: None
* *
* *
     Rechallenge: None
* *
* *
     Thirty guinea pigs were allocated to two groups: a control group 1 (five
     males and five females) and a treated group 2 (ten males and ten
     females).
* *
     Day 1: Intradermal injections of Freund's complete
F007 Clinical Signs: None
* *
     Rechallenge: None
* *
* *
     Thirty guinea pigs were allocated to two groups: a control group 1 (five
     males and five females) and a treated group 2 (ten males and ten
     females).
* *
     Day 1: Intradermal injections of Freund's complete adjuvant mixed with
     the test substance (treated group) or the vehicle (control group) were
     performed in the dorsal region between the shoulders. Day 7: The same
     dorsal region between the shoulders received a topical application of
     sodium lauryl sulfate in Vaseline (10% w/w) in order to induce local
     irritation.
* *
     Day 8: The test site was treated by topical application of the test
     substance (treated group) or the vehicle (control group) and was covered
     by an occlusive dressing for 48 hours.
* *
     Day 22: After a rest period of 12 days, all animals of the treated and
     control groups were challenged by a topical application of the test
     substance to the right flank. The left flank served as control and
     received the vehicle only. Test substance and vehicle were maintained
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under an occlusive dressing for 24 hours. Skin reactions were evaluated
     approximately 24 and 48 hours later.
* *
* *
     At the end of the study, animals were killed without examination of
     internal organs. No skin samples were taken from the challenge
     application sites.
* *
     The sensitivity of the guinea pigs was checked with a positive
     sensitizer: 2,4-dinitro chlorobenzene (DNCB). During the induction
     period, the test substance was applied at 0.1 % (w/w) (day 1) and 1 %
     (w/w) (day 8). For the challenge application, the DNCB was applied to the
     right flank at a concentration of 0.5\% (w/w).
F008 CLGETTS
F012 20
F020 120238
EOR
F002 27
F010 5.3
F004 2
F005 RS
F006 sBA is not a dermal sensitizer in guinea pigs. The sensitivity of the
     guinea pigs was satisfactory since 50% of the animals showed a positive
     reaction with positive control sensitizer, DNCB. Scores in control and
     treated groups after challe
F007 sBA is not a dermal sensitizer in guinea pigs. The sensitivity of the
     guinea pigs was satisfactory since 50% of the animals showed a positive
     reaction with positive control sensitizer, DNCB. Scores in control and
     treated groups after challenge with sBA were 0 at both 24 and 48 hours.
F008 CLGETTS
F012 20
F020 120239
EOR
F002 27
F010 5.3
F004 2
F005 SO
F006 Centre d'Elevage Lebeau, Gambais, France
F007 Centre d'Elevage Lebeau, Gambais, France
F008 CLGETTS
F012 20
F020 120240
EOR
F002 27
F010 5.3
F004 2
F005 TS
F006 sec-Butanol (sBA) 99.87%; ATOFINA Chemicals, Inc,
F007 sec-Butanol (sBA) 99.87%; ATOFINA Chemicals, Inc,
F008 CLGETTS
F012 20
F020 120241
EOR
F002 27
F010 5.4
F004 1
F005 CL
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F006 MEK has a low order of toxicity based on the 90-day repeated-dose

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exposures. The NOAEL is 2500 ppm.
F007 MEK has a low order of toxicity based on the 90-day repeated-dose
     exposures. The NOAEL is 2500 ppm.
F008 CLGETTS
F012 20
F020 120242
EOR
F002 27
F010 5.4
F004 1
F005 RE
F006 Cavender, F.L., Casey, H.W., Salem, H., Swenberg, J.A., and Gralla, E.J.
     (1983). A 90-day vapor inhalation toxicity study of methyl ethyl ketone.
     Fund. Appl. Toxicol. 3:264-270.
F007 Cavender, F.L., Casey, H.W., Salem, H., Swenberg, J.A., and Gralla, E.J.
     (1983). A 90-day vapor inhalation toxicity study of methyl ethyl ketone.
      Fund. Appl. Toxicol. 3:264-270.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120243
EOR
F002 27
F010 5.4
F004 1
F005 RE
F006 Toxigenics. (1981). 90-Day vapor inhalation study of methyl ethyl ketone
     in albino rats. Toxigenics' Study 420-0305. Toxigenics, Inc., 1800 East
     Pershing Road, Decatur, IL 62526.
F007 Toxigenics. (1981). 90-Day vapor inhalation study of methyl ethyl ketone
     in albino rats. Toxigenics' Study 420-0305. Toxigenics, Inc., 1800 East
     Pershing Road, Decatur, IL 62526.
F008 HEDSET
F012 20
F020 120244
EOR
F002 27
F010 5.4
F004 1
F005 RL
F006 1 - Reliable study with restrictions. Study well documented, meets
     generally accepted scientific principles, acceptable for assessment.
F007 1 - Reliable study with restrictions. Study well documented, meets
     generally accepted scientific principles, acceptable for assessment.
F008 HEDSET
F012 20
F020 120245
EOR
F002 27
F010 5.4
F004 1
F005 RM
F006 An extensive pathologic investigation was conducted and no lesions were
     found that could be attributed to MEK exposure. There was no indication
     that repeated exposure to relatively high levels of MEK had any effect on
     reproductive tissues.
F007 An extensive pathologic investigation was conducted and no lesions were
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found that could be attributed to MEK exposure. There was no indication
     that repeated exposure to relatively high levels of MEK had any effect on
     reproductive tissues. The examined tissues included testes,
     epididymides, seminal vesicles, vagina, cervix, uterus, oviducts, and
     ovaries.
F008 HEDSET
F012 20
F020 120246
EOR
F002 27
F010 5.4
F004 1
F005 RM
F006 Methyl ethyl ketone is metabolically interchangeable with
     2-butanol. Approximately 97% of a 2-butanol dose is
     oxidized to methyl ethyl ketone.
F007 Methyl ethyl ketone is metabolically interchangeable with
     2-butanol. Approximately 97% of a 2-butanol dose is
* *
     oxidized to methyl ethyl ketone.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120247
EOR
F002 27
F010 5.4
F004 1
F005 RM
F006 Purity: > 99.5%
    Number/Sex/Dose: 15 (10/sex - principals - for routine pathology and
     5/sex - dedicated - for special neuropathology)
    Vehicle: None
F007 Purity: > 99.5%
    Number/Sex/Dose: 15 (10/sex - principals - for routine pathology and
     5/sex - dedicated - for special neuropathology)
     Vehicle: None
F008 HEDSET
F012 20
F020 120248
EOR
F002 27
F010 5.4
F004 1
F005 RS
F006 The 90-day exposures had no adverse effect on the clinical health or
     growth of male or female rats except for a depression of mean body weight
     in the 5000-ppm exposure group. No animals died during the study. No
     signs of nasal irritation
F007 The 90-day exposures had no adverse effect on the clinical health or
     growth of male or female rats except for a depression of mean body weight
     in the 5000-ppm exposure group. No animals died during the study.
     signs of nasal irritation were observed during the study. There were no
     treatment-related effects in food consumption or ophthalmologic studies
     in any of the rats exposed to MEK vapors. The 5000-ppm animals had a
     slight but significant increase in liver weight, liver weight/body weight
    ratio and liver weight/brain weight ratio at the time of necropsy. Serum
     glutamic-pyruvic transaminase (SGPT) activity in the 2500-ppm female rats
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was elevated while the 5000-ppm female rats exhibited significantly
     decreased SGPT activity. In addition, alkaline phosphatase, potassium,
     and glucose values for the 5000-ppm female rats were increased. Special
     neuropathological and routine pathological studies did not reveal any
     lesions that could be attributed to MEK exposure.
F008 HEDSET
F012 20
F020 120249
EOR
F002 27
F010 5.4
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120250
EOR
F002 27
F010 5.4
F004 1
F005 TS
F006 Methyl Ethyl Ketone (MEK)
F007 Methyl Ethyl Ketone (MEK)
F008 CLGETTS
F012 20
F020 120251
EOR
F002 27
F010 5.4
F004 2
F005 CL
F006 sBA (probably through its metabolite methyl-ethyl-ketone) was a potent
     inducer of P-450 in the kidney and liver.
F007 sBA (probably through its metabolite methyl-ethyl-ketone) was a potent
     inducer of P-450 in the kidney and liver.
F008 CLGETTS
F012 20
F020 120252
EOR
F002 27
F010 5.4
F004 2
F006 Aarstad K., Zahlsen, K., and Nilsen, O. G. (1985). Inhalation of
    butanols: changes in the cytochrome p-450 enzyme system. Arch Toxycol
     (Suppl) 8:418-421.
F007 Aarstad K., Zahlsen, K., and Nilsen, O. G. (1985). Inhalation of
    butanols: changes in the cytochrome p-450 enzyme system. Arch Toxycol
     (Suppl) 8:418-421.
F008 CLGETTS
F012 20
F020 120253
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EOR
F002 27
F010 5.4
F004 2
F005 RL
F006 2- Reliable study with restrictions. Study well documented, meets
     generally accepted scientific principles, acceptable for assessment.
F007 2- Reliable study with restrictions. Study well documented, meets
     generally accepted scientific principles, acceptable for assessment.
F008 CLGETTS
F012 20
F020 120254
EOR
F002 27
F010 5.4
F004 2
F005 RM
F006 Purity: > or = 99%
   Number/Sex/Dose: 5
* *
    Vehicle: None
F007 Purity: > or = 99%
** Number/Sex/Dose: 5
* *
    Vehicle: None
F008 CLGETTS
F012 20
F020 120255
EOR
F002 27
F010 5.4
F004 2
F005 RS
F006 sBA caused increases in cytochrome P-450 concentrations in the kidneys
     (47% rise; 500 ppm for 5 days) and liver (33% rise; 2000 ppm for 3 days).
     Slight decreases in lung cytochrome P-450 were observed.
F007 sBA caused increases in cytochrome P-450 concentrations in the kidneys
     (47% rise; 500 ppm for 5 days) and liver (33% rise; 2000 ppm for 3 days).
     Slight decreases in lung cytochrome P-450 were observed.
F008 CLGETTS
F012 20
F020 120256
EOR
F002 27
F010 5.5
F004 1
F005 CL
F006 Not Mutagenic
F007 Not Mutagenic
F008 CLGETTS
F012 20
F020 120257
EOR
F002 27
F010 5.5
F004 1
F005 RE
F006 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
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3(3):227-231.
F007 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
     3(3):227-231.
F008 CLGETTS
F012 20
F020 120258
EOR
F002 27
F010 5.5
F004 1
F005 RL
F006 1- Reliable without restriction
F007 1- Reliable without restriction
F008 CLGETTS
F012 20
F020 120259
EOR
F002 27
F010 5.5
F004 1
F005 RM
F006 Purity: 99.5%
     Strain: S. typhimurium / TA98, TA100, TA1535, TA1537, TA1538
     Species/Cell Type: Rat Liver (S9 fraction)
* *
    Vehicle: DMSO
F007 Purity: 99.5%
     Strain: S. typhimurium / TA98, TA100, TA1535, TA1537, TA1538
     Species/Cell Type: Rat Liver (S9 fraction)
    Vehicle: DMSO
F008 CLGETTS
F012 20
F020 120260
EOR
F002 27
F010 5.5
F004 1
F005 RM
F006 sBA did not induce reverse gene mutation in bacteria.
F007 sBA did not induce reverse gene mutation in bacteria.
F008 CLGETTS
F012 20
F020 120261
EOR
F002 27
F010 5.5
F004 1
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120262
EOR
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F002 27
F010 5.5
F004 1
F005 TC
F006 Control plates were set up with solvent alone and with an appropriate
    known positive control compound. All tests were carried out in
     triplicate. Two replicate assays were carried on different days in order
     to confirm the reproducibility o
F007 Control plates were set up with solvent alone and with an appropriate
    known positive control compound. All tests were carried out in
     triplicate. Two replicate assays were carried on different days in order
     to confirm the reproducibility of the results. The S9 fractions were
    prepared from Aroclor-induced rats. Initially, a range of test
    concentrations of test material was tested and a second experiment was
    then performed based on the results taking into account the effect on
     cell viability and any possible positive increases in mitotic gene
     conversion. Control plates were set up with solvent alone and with the
    positive control compounds 4-nitroquinoline-N-oxide and cyclophosphamide.
F008 CLGETTS
F012 20
F020 120263
EOR
F002 27
F010 5.5
F004 2
F005 CL
F006 Not Mutagenic
F007 Not Mutagenic
F008 CLGETTS
F012 20
F020 120264
EOR
F002 27
F010 5.5
F004 2
F005 RE
F006 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
     3(3):227-231.
F007 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
     3(3):227-231.
F008 CLGETTS
F012 20
F020 120265
EOR
F002 27
F010 5.5
F004 2
F005 RL
F006 1- Reliable without restriction.
F007 1- Reliable without restriction.
F008 CLGETTS
F012 20
F020 120266
F:OR
F002 27
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F010 5.5
F004 2
F005 RM
F006 Purity: 99.5%
     Strain: E. coli WP2 uvr A
* *
     Species/Cell Type: Rat Liver (S9 fraction)
* *
    Vehicle: DMSO
F007 Purity: 99.5%
     Strain: E. coli WP2 uvr A
* *
     Species/Cell Type: Rat Liver (S9 fraction)
    Vehicle: DMSO
F008 CLGETTS
F012 20
F020 120267
EOR
F002 27
F010 5.5
F004 2
F005 RM
F006 sBA did not induce reverse gene mutation in bacteria.
F007 sBA did not induce reverse gene mutation in bacteria.
F008 CLGETTS
F012 20
F020 120268
EOR
F002 27
F010 5.5
F004 2
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120269
EOR
F002 27
F010 5.5
F004 2
F005 TC
F006 Control plates were set up with solvent alone and with an appropriate
     known positive control compound. All tests were carried out in
     triplicate. Two replicate assays were carried on different days in order
     to confirm the reproducibility o
F007 Control plates were set up with solvent alone and with an appropriate
    known positive control compound. All tests were carried out in
     triplicate. Two replicate assays were carried on different days in order
     to confirm the reproducibility of the results. The S9 fractions were
    prepared from Aroclor-induced rats. Initially, a range of test
     concentrations of test material was tested and a second experiment was
     then performed based on the results taking into account the effect on
     cell viability and any possible positive increases in mitotic gene
     conversion. Control plates were set up with solvent alone and with the
     positive control compounds 4-nitroquinoline-N-oxide and cyclophosphamide.
F008 CLGETTS
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F012 20
F020 120270
EOR
F002 27
F010 5.5
F004 3
F005 CL
F006 Not mutagenic
F007 Not mutagenic
F008 CLGETTS
F012 20
F020 120271
EOR
F002 27
F010 5.5
F004 3
F005 RE
F006 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
     3(3):227-231.
F007 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
     3(3):227-231.
F008 CLGETTS
F012 20
F020 120272
EOR
F002 27
F010 5.5
F004 3
F005 RL
F006 1- Reliable without restriction
F007 1- Reliable without restriction
F008 CLGETTS
F012 20
F020 120273
EOR
F002 27
F010 5.5
F004 3
F005 RM
F006 Purity: 99.5%
     Strain: Saccharomyces cerevisiae
     Species/Cell Type: Rat Liver (S9 fraction)
* *
    Vehicle: DMSO
F007 Purity: 99.5%
* *
     Strain: Saccharomyces cerevisiae
     Species/Cell Type: Rat Liver (S9 fraction)
    Vehicle: DMSO
F008 CLGETTS
F012 20
F020 120274
EOR
F002 27
F010 5.5
F004 3
F005 RM
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F006 sBA did not induce mitotic gene conversion in yeast.
F007 sBA did not induce mitotic gene conversion in yeast.
F008 CLGETTS
F012 20
F020 120275
EOR
F002 27
F010 5.5
F004 3
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120276
EOR
F002 27
F010 5.5
F004 3
F005 TC
F006 Yeast cells were grown in log phase, washed and re-suspended in 2/5
     strength YEPD broth at a concentration of 10 X 106cells/ml. The
     suspension was then divided into 1.9 ml amounts in 30-ml universal
     containers and 0.1 ml of the test compou
F007 Yeast cells were grown in log phase, washed and re-suspended in 2/5
     strength YEPD broth at a concentration of 10 X 106cells/ml. The
     suspension was then divided into 1.9 ml amounts in 30-ml universal
     containers and 0.1 ml of the test compound solution was added (-S9). For
     the experiments with metabolic activation (+S9), 0.1 ml of the compound
     was added to 1.6 ml of yeast suspension, together with 0.3 ml of S9 mix.
     The cultures were incubated with shaking, with shaking, at 30°C for 18
     hours. Aliquots were then plated onto the appropriate culture media for
     the selection of mitotic gene convertants and cells surviving the
     treatment.
F008 CLGETTS
F012 20
F020 120277
EOR
F002 27
F010 5.5
F004 4
F005 CL
F006 Not Mutagenic
F007 Not Mutagenic
F008 CLGETTS
F012 20
F020 120278
EOR
F002 27
F010 5.5
F004 4
F005 RE
F006 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
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3(3):227-231.
F007 Brooks, T. M., Meyer, A., and Hutson, D. H. (1988). The genetic
     toxicology of some hydrocarbon and oxygenated solvents. Mutagenesis,
     3(3):227-231.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120279
EOR
F002 27
F010 5.5
F004 4
F005 RL
F006 1- Reliable without restriction
F007 1- Reliable without restriction
F008 HEDSET
F012 20
F020 120280
EOR
F002 27
F010 5.5
F004 4
F005 RM
F006 Purity: 99.5%
    Species/Cell Type: Chinese Hamster Ovary Cells
    Vehicle: DMSO
F007 Purity: 99.5%
     Species/Cell Type: Chinese Hamster Ovary Cells
     Vehicle: DMSO
F008 HEDSET
F012 20
F020 120281
EOR
F002 27
F010 5.5
F004 4
F005 RM
F006 sBA did not induce chromosome damage in CHO mammalian cells.
F007 sBA did not induce chromosome damage in CHO mammalian cells.
F008 HEDSET
F012 20
F020 120282
EOR
F002 27
F010 5.5
F004 4
F005 RS
F006 Negative - No increase in chromosomal aberrations
F007 Negative - No increase in chromosomal aberrations
F008 HEDSET
F012 20
F020 120283
EOR
F002 27
F010 5.5
F004 4
F005 S0
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F006 EXXON CHEMICAL, Limited Fareham, Hampshire
** EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120284
EOR
F002 27
F010 5.5
F004 4
F005 TC
F006 Two separate cytotoxicity assays were performed: (i) Monolayer CHO
     cultures were prepared in multi-well tissue culture trays. The cultures
     were incubated at 37°C for 24 hours to commence active growth before
     treatment with the test materia
F007 Two separate cytotoxicity assays were performed: (i) Monolayer CHO
     cultures were prepared in multi-well tissue culture trays. The cultures
     were incubated at 37°C for 24 hours to commence active growth before
     treatment with the test material. Assays were performed both in the
    presence and in the absence of S9 mix using either a 5-hour exposure
     (+S9) or a 24-hour exposure (-S9). Twenty-four hours after initial
    treatment, cultures were stained with Giemsa and the growth inhibition
    was noted. (ii) CHO cultures were prepared in 25 cm2 flasks, and after
     24 hours the cells were exposed to sBA both in the presence and in the
    absence of S9 mix. The number of cell in each flask was counted 24 hours
     later. The compound concentrations selected for the chromosome assays
    were 1, 0.5, and 0.25 times the GI50 level.
* *
    CHO Chromosome Assay: Cultured CHO cells were grown in 80-cm2 flasks for
     24 hours before treatment. Treatment periods were 5hours in the presence
    of S9 mix and 24 hours in the absence of S9 mix. Positive control
    cultures were run in parallel [ethyl methanesulphonate (-S9) and
    cyclophosphamide (+S9). Colcemid was added to all cultures 22 hours after
    treatment. After a further 2 hours, the cells were trypsinized,
    resuspended in hypotonic solution and then fixative, before spotting onto
     slides. Cell preparations were then stained with Giemsa. The slides
    were randomly coded and 100 cells from each culture were analyzed
    microscopically. Mitotic index estimations were also made.
F008 CLGETTS
F012 20
F020 120285
EOR
F002 27
F010 5.5
F004 5
F005 CL
F006 Not Mutagenic
F007 Not Mutagenic
F008 CLGETTS
F012 20
F020 120286
EOR
F002 27
F010 5.5
F004 5
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F005 RE
F006 ELF ATOCHEM, BUTANOL SECONDAIRE TEST DE AMES "essai de mutation reverse
     sur Salmonella typhirium" SANOFI RECHERCHE Report CEL357A/C, 1989.
F007 ELF ATOCHEM, BUTANOL SECONDAIRE TEST DE AMES "essai de mutation reverse
     sur Salmonella typhirium" SANOFI RECHERCHE Report CEL357A/C, 1989.
F008 HEDSET
F009 02-06-1994
F012 20
F020 120287
EOR
F002 27
F010 5.5
F004 5
F005 RL
F006 1- Reliable without restriction
F007 1- Reliable without restriction
F008 HEDSET
F012 20
F020 120288
EOR
F002 27
F010 5.5
F004 5
F005 RM
F006 sBA did not induce reverse gene mutation in bacteria.
F007 sBA did not induce reverse gene mutation in bacteria.
F008 HEDSET
F012 20
F020 120289
EOR
F002 27
F010 5.5
F004 5
F005 RM
F006 Strain: Salmonella typhimurium / TA98, TA100, TA1535, TA1537, TA1538
     Species/Cell Type: Rat Liver (S9 fraction)
     Vehicle: DMSO
F007 Strain: Salmonella typhimurium / TA98, TA100, TA1535, TA1537, TA1538
     Species/Cell Type: Rat Liver (S9 fraction)
    Vehicle: DMSO
F008 HEDSET
F012 20
F020 120290
EOR
F002 27
F010 5.5
F004 5
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120291
EOR
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```
F002 27
F010 5.5
F004 5
F005 TC
F006 Control plates were set up with solvent alone and with an appropriate
     known positive control compound. All tests were carried out in
     triplicate. Two replicate assays were carried out on different days in
     order to confirm the reproducibili
F007 Control plates were set up with solvent alone and with an appropriate
    known positive control compound. All tests were carried out in
     triplicate. Two replicate assays were carried out on different days in
     order to confirm the reproducibility of the results. The S9 fractions
    were prepared from Aroclor -induced rats. Initially, a range of test
     concentrations of test material was tested and a second experiment was
     then performed based on the results taking into account the effect on
     cell viability and any possible positive increases in mitotic gene
     conversion. Control plates were set up with solvent alone and with the
     positive control compounds 4-nitroquinoline-N-oxide and cyclophosphamide.
F008 CLGETTS
F012 20
F020 120292
EOR
F002 27
F010 5.6
F004 1
F005 CL
F006 unknown
F007 unknown
F008 CLGETTS
F012 20
F020 120293
EOR
F002 27
F010 5.6
F004 1
F005 RE
F006 Barilyak, I. R. and Kozachuk, SYu. (1988). Investigation of the
     cytogenetic effect of a number of monohydric alcohols on rat bone marrow
     cells. Cytology and Genetics (Tsitologiaya I Genetika), 22(2):49-52.
F007 Barilyak, I. R. and Kozachuk, SYu. (1988). Investigation of the
     cytogenetic effect of a number of monohydric alcohols on rat bone marrow
     cells. Cytology and Genetics (Tsitologiaya I Genetika), 22(2):49-52.
F008 CLGETTS
F012 20
F020 120294
EOR
F002 27
F010 5.6
F004 1
F005 RL
F006 4 - Incomplete translation from Russian and insufficient for assessment
F007 4 - Incomplete translation from Russian and insufficient for assessment
F008 CLGETTS
F012 20
F020 120295
EOR
F002 27
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F010 5.6
F004 1
F005 RM
F006 Single intragastric administration of monohydric alcohols at equitoxic
     doses (1/5 LD50) affected the chromosomal appearance of bone marrow.
F007 Single intragastric administration of monohydric alcohols at equitoxic
     doses (1/5 LD50) affected the chromosomal appearance of bone marrow.
F008 CLGETTS
F012 20
F020 120296
EOR
F002 27
F010 5.6
F004 1
F005 RM
F006 System of Testing: Rat Bone Marrow
    Metabolic Activation: unknown
    Species/Cell Type: Rat
* *
    Concentrations Tested: 1/5 of the LD50 by intragastric route
* *
    Vehicle: unknown
F007 System of Testing: Rat Bone Marrow
    Metabolic Activation: unknown
* *
    Species/Cell Type: Rat
* *
    Concentrations Tested: 1/5 of the LD50 by intragastric route
    Vehicle: unknown
F008 CLGETTS
F012 20
F020 120297
EOR
F002 27
F010 5.6
F004 1
F005 RS
F006 Increased numbers of polyploid cells, cells with chromosome gaps, and
     cells with chromosomal aberrations.
F007 Increased numbers of polyploid cells, cells with chromosome gaps, and
     cells with chromosomal aberrations.
F008 CLGETTS
F012 20
F020 120298
EOR
F002 27
F010 5.6
F004 2
F005 CL
F006 MEK did not induce a statistically significant increase in the mean
     number of micronucleated polychromatic erythrocytes in the bone marrow of
     CD-1 mice. Therefore, it is not considered mutagenic under the
     conditions of this assay.
F007 MEK did not induce a statistically significant increase in the mean
     number of micronucleated polychromatic erythrocytes in the bone marrow of
     CD-1 mice. Therefore, it is not considered mutagenic under the
     conditions of this assay.
F008 CLGETTS
F012 20
F020 120299
EOR
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```
F002 27
F010 5.6
F004 2
F005 RE
F006 O'Donoghue, J.L., Haworth, S.R., Curren, R.D., Kirby, P.E., Lawlor, T.,
     Moran, E.J., Phillips, R.D., Putnam, D.L., Rogers-Back, A.M., Slesinski,
     R.S., and Thilagar, A. (1988). Mutagenicity studies on ketone solvents:
     methyl ethyl ketone, me
F007 O'Donoghue, J.L., Haworth, S.R., Curren, R.D., Kirby, P.E., Lawlor, T.,
     Moran, E.J., Phillips, R.D., Putnam, D.L., Rogers-Back, A.M., Slesinski,
     R.S., and Thilagar, A. (1988). Mutagenicity studies on ketone solvents:
     methyl ethyl ketone, methyl isobutyl ketone, and isophorone. Mutation
     Research, 206:149-161.
F008 CLGETTS
F012 20
F020 120300
EOR
F002 27
F010 5.6
F004 2
F005 RL
F006 1 - Reliable without Restrictions. No circumstances occurred that would
    have affected the quality or integrity of the data.
F007 1 - Reliable without Restrictions. No circumstances occurred that would
    have affected the quality or integrity of the data.
F008 CLGETTS
F012 20
F020 120301
EOR
F002 27
F010 5.6
F004 2
F005 RM
F006 Not mutagenic
F007 Not mutagenic
F008 CLGETTS
F012 20
F020 120302
EOR
F002 27
F010 5.6
F004 2
F005 RM
F006 Purity: 99.9%
     Number: 5/sex/dose
     Dose/Time: 1.96 ml/kg Single injection (dose equal to the MEK LD20
     reported as ml test article / kg body weight when administered in a total
     volume of 10 ml test article-vehicle mixture / kg body weight).
F007 Purity: 99.9%
     Number: 5/sex/dose
* *
     Dose/Time: 1.96 ml/kg Single injection (dose equal to the MEK LD20
     reported as ml test article / kg body weight when administered in a total
     volume of 10 ml test article-vehicle mixture / kg body weight).
* *
     Vehicle: Corn Oil (10 ml/kg)
* *
     Positive Control: 0.25 mg/kg Triethylene melamine (TEM)
F008 CLGETTS
F012 20
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```
F020 120303
EOR
F002 27
F010 5.6
F004 2
F005 RM
F006 The test substance and the vehicle were administered as a single dose by
     intraperitoneal injection. The vehicle was dosed at a volume equal to
     the test substance volume. The positive control was administered as a
     single dose at a volume e
F007 The test substance and the vehicle were administered as a single dose by
     intraperitoneal injection. The vehicle was dosed at a volume equal to
     the test substance volume. The positive control was administered as a
     single dose at a volume equal to the test substance volume. Animals from
     the appropriate groups were sacrificed at approximately 12, 24 and 48
    hours. Animals dosed with triethylene melamine were sacrificed at 24
    hours only. Immediately following sacrifice, the femur was exposed and
    the bone marrow was aspirated into a syringe containing fetal calf serum.
     The cells were washed, centrifuged, and resuspended. Slide smears of
     the bone marrow were made for each animal and stained with
    May-Gruenwald-Giemsa stain. Coded slides were then evaluated for
    presence of micronuclei (1000 polychromatic erythrocytes/animal were
    evaluated). A 1-way analysis of variance and Duncan's multiple range
     test (p = 0.05) were used to assess the statistical significance of any
     observed effects.
F008 CLGETTS
F012 20
F020 120304
EOR
F002 27
F010 5.6
F004 2
F005 RS
F006 None of the dose groups were statistically different from the vehicle
     control. The positive control (0.25 mg/kg TEM) induced a statistically
     significant increase in the mean number of micronucleated polychromatic
     erythrocytes (p = 0.01) wh
F007 None of the dose groups were statistically different from the vehicle
     control. The positive control (0.25 mg/kg TEM) induced a statistically
     significant increase in the mean number of micronucleated polychromatic
     erythrocytes (p = 0.01) which indicates that the positive control was
    clastogenic and the test system responded in an appropriate manner.
    Vehicle carrier control values for the mean percent of polychromatic
     erythrocytes and for the mean percent of micronucleated polychromatic
     erythrocytes responded in an appropriate manner.
F008 CLGETTS
F012 20
F020 120305
EOR
F002 27
F010 5.6
F004 2
F005 TS
F006 Methyl Ethyl Ketone (MEK) CAS No. 78-93-3
F007 Methyl Ethyl Ketone (MEK) CAS No. 78-93-3
F008 CLGETTS
F012 20
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F020 120306
EOR
F002 27
F010 5.7
F004 1
F005 RM
F006 No studies of 2-butanol for carcinogenic activity have been
     reported; however, based on results of mutagenicity assays
     it is expected to have a low potential for carcinogenicity.
F007 No studies of 2-butanol for carcinogenic activity have been
     reported; however, based on results of mutagenicity assays
     it is expected to have a low potential for carcinogenicity.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120307
EOR
F002 27
F010 5.7
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120308
EOR
F002 27
F010 5.8.1
F004 1
F005 CL
F006 sBA is not a reproductive hazard.
F007 sBA is not a reproductive hazard.
F008 CLGETTS
F012 20
F020 120309
EOR
F002 27
F010 5.8.1
F004 1
F005 RE
F006 Cox, G.E., D.E. Bailey and K. Morgareidge. (1975). Toxicity studies in
     rats with 2-butanol including growth, reproduction and teratologic
     observations. LaB No. 2093. Food and Drug Research Laboratories, Inc.,
     Waverly, NY.
F007 Cox, G.E., D.E. Bailey and K. Morgareidge. (1975). Toxicity studies in
     rats with 2-butanol including growth, reproduction and teratologic
     observations. LaB No. 2093. Food and Drug Research Laboratories, Inc.,
     Waverly, NY.
F008 HEDSET
F009 31-03-1994
F012 20
F020 120310
EOR
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```
F002 27
F010 5.8.1
F004 1
F005 RE
F006 Gallo, M.A., Oser, B.L., Cox, G.E., and Bailey, D.E. (1977). Studies on
     the long-term toxicity of 2-butanol. Toxicology and Applied Pharmacology,
     41:135 (Abstract).
F007 Gallo, M.A., Oser, B.L., Cox, G.E., and Bailey, D.E. (1977). Studies on
     the long-term toxicity of 2-butanol. Toxicology and Applied Pharmacology,
     41:135 (Abstract).
F008 HEDSET
F012 20
F020 120311
EOR
F002 27
F010 5.8.1
F004 1
F005 RL
F006 2 - Reliable study with restrictions. No circumstances occurred that
     would have affected the quality or integrity of the data. Comparable to a
     guideline study and test procedures were in accordance with generally
     accepted scientific standa
F007 2 - Reliable study with restrictions. No circumstances occurred that
     would have affected the quality or integrity of the data. Comparable to a
     guideline study and test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
F008 HEDSET
F012 20
F020 120312
EOR
F002 27
F010 5.8.1
F004 1
F005 RM
F006 Purity: > or = 99%
* *
     Number/Sex/Dose: 30
* *
     Doses/Concentration:
* *
     F0 Generation: 0, 0.3, 1.0, or 3.0% solutions (0, 538, 1644, and 5089
     mg/kg-day for males and 0, 594, 1771, and 4571 mg/kg-day for females)
* *
     F1 Generation: 0, 0.3, 1.0, or 2.0%
F007 Purity: > or = 99%
* *
* *
     Number/Sex/Dose: 30
* *
* *
     Doses/Concentration:
* *
     F0 Generation: 0, 0.3, 1.0, or 3.0% solutions (0, 538, 1644, and 5089
     mg/kg-day for males and 0, 594, 1771, and 4571 mg/kg-day for females)
* *
     F1 Generation: 0, 0.3, 1.0, or 2.0% (2.0% calculated to be equivalent to
     3384 mg/kg-day for males and 3122 mg/kg-day for females)
* *
     Vehicle: Drinking Water
F008 HEDSET
F012 20
F020 120313
EOR
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F002 27
F010 5.8.1
F004 1
F005 RM
F006 sec-Butanol was initially administered to the F0 generation at
     concentrations of 0, 0.3, 1.0, and 3.0% in the drinking water. Due to
     toxicity, the high level was reduced to 2.0% during the second-generation
     study (F1). The F1 generation a
F007 sec-Butanol was initially administered to the F0 generation at
     concentrations of 0, 0.3, 1.0, and 3.0% in the drinking water.
     toxicity, the high level was reduced to 2.0% during the second-generation
     study (F1). The F1 generation animals (30/sex/group) were reared to
    maturity (up to week 12), mated to produce a F2 generation, then
     sacrificed for organ weights, and gross and microscopic pathological
     evaluations (10/sex/group). Hematological, biochemical, and urinary
     examinations were conducted terminally on the F1 rats. A series of mild
    changes in the kidney (non-reactive tubular degeneration, tubular casts,
    foci of tubular regeneration, microcysts) were observed animals treated
    at 2.0% sBA. The authors concluded that these findings were not a result
    of direct toxicity and did not have clear pathologic significance.
    Rather they were non-specific effects due to increased renal work load,
    possibly from increased urine volume and pressure at the high dose of sBA
     (Cox et al, 1975). No other findings of note were seen. The no-effect
     level for the study was 1.0% (estimated to be 1500 mg/kg/day by the
    authors and 1771 mg/kg/day by EPA/IRIS).
F008 HEDSET
F012 20
F020 120314
EOR
F002 27
F010 5.8.1
F004 1
F005 RS
F006 Maternal NOEL: 1% (~1500 mg/kg/day)
    Maternal NOAEL: 1771 mg/kg/day
     Pup NOEL: 1% (~1500 mg/kg/day)
* *
    Pup NOAEL: 1771 mg/kg/day
F007 Maternal NOEL: 1% (~1500 mg/kg/day)
    Maternal NOAEL: 1771 mg/kg/day
* *
    Pup NOEL: 1% (~1500 mg/kg/day)
    Pup NOAEL: 1771 mg/kg/day
F008 HEDSET
F012 20
F020 120315
EOR
F002 27
F010 5.8.1
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
    EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120316
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EOR
F002 27
F010 5.8.2
F004 1
F005 CL
F006 sBA is not a teratogen. There was no evidence of teratogenic events nor
    was there evidence of selective developmental toxicity.
F007 sBA is not a teratogen. There was no evidence of teratogenic events nor
    was there evidence of selective developmental toxicity.
F008 CLGETTS
F012 20
F020 120317
EOR
F002 27
F010 5.8.2
F004 1
F005 ME
F006 For the maternal data, multivariate analysis (with baseline as covariant)
     was used for weight comparisons across groups. The group differences in
     food and water intake were analyzed by multivariate analysis of variance.
     A Kruskal-Wallis te
F007 For the maternal data, multivariate analysis (with baseline as covariant)
    was used for weight comparisons across groups. The group differences in
     food and water intake were analyzed by multivariate analysis of variance.
     A Kruskal-Wallis test was used for group comparisons of corpora lutea
    per animal. For the fetal data, analysis of covariance was used to
    compare fetal weights across groups and sex. Group comparisons of the
    variables including litter size, percentage alive/litter, percentage
    normal/litter, and percentage females/litter were made using
    Kruskal-Wallis test. For the variables including skeletal malformations,
     skeletal variations, visceral malformations, visceral variations,
    external malformations, and non-normal fetuses, the number of litters
    with one or more of the variables of interest was compared between groups
    using Fisher's exact test. The results of the test were adjusted for
    multiple comparisons, when appropriate, using the Bonferroni technique.
     A probability of p = < 0.05 was required for significance.
F008 CLGETTS
F012 20
F020 120318
EOR
F002 27
F010 5.8.2
F004 1
F005 RE
F006 Nelson, B. K., Brightwell, W. S., Khan, A., Burg, J. R., and Goad, P. T.
     (1989). Lack of selective developmental toxicity of three butanol
     isomers administered by inhalation to rats. Fundam Appl Toxicol,
     12:469-479.
F007 Nelson, B. K., Brightwell, W. S., Khan, A., Burg, J. R., and Goad, P. T.
     (1989). Lack of selective developmental toxicity of three butanol
     isomers administered by inhalation to rats. Fundam Appl Toxicol,
     12:469-479.
F008 HEDSET
F009 21-10-1992
F012 20
F020 120319
EOR
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F002 27
F010 5.8.2
F004 1
F005 RL
F006 2- Reliable study with restrictions. Study well documented, meets
    generally accepted scientific principles, acceptable for assessment.
F007 2- Reliable study with restrictions. Study well documented, meets
    generally accepted scientific principles, acceptable for assessment.
F008 HEDSET
F012 20
F020 120320
EOR
F002 27
F010 5.8.2
F004 1
F005 RM
F006 Groups of 15-16 rats were exposed by inhalation to 0, 3,500, 5,000 or
     7,000 ppm sBA, 7 hours/day on days 1-19 of gestation; the dams were
     sacrificed on day 20. At 7,000 ppm, narcosis was observed in all
     animals. At 5000 ppm, the dams were
F007 Groups of 15-16 rats were exposed by inhalation to 0, 3,500, 5,000 or
     7,000 ppm sBA, 7 hours/day on days 1-19 of gestation; the dams were
     sacrificed on day 20. At 7,000 ppm, narcosis was observed in all
     animals. At 5000 ppm, the dams were partially narcotized with locomotion
    activity impaired. Maternal weight gain and food consumption were
     significantly reduced in all dose groups. No data collected on maternal
    organ weights, or gross or microscopic lesions. The number of live
     fetuses was significantly reduced and resorptions were increased in the
    high exposure group only. Fetal body weights were significantly reduced
     in the mid- and high dose groups. There was no evidence of teratogenic
     effects in this study, and there was also no evidence of selective
     developmental toxicity. The no-effect levels were < 3,500 ppm for
    maternal toxicity and 3,500 ppm for developmental toxicity.
F008 HEDSET
F012 20
F020 120321
EOR
F002 27
F010 5.8.2
F004 1
F005 RM
F006 Purity: > or = 99%
    Number/Sex/Dose: 15-16 Mated Females
    Vehicle: None
F007 Purity: > or = 99%
    Number/Sex/Dose: 15-16 Mated Females
* *
    Vehicle: None
F008 HEDSET
F012 20
F020 120322
EOR
F002 27
F010 5.8.2
F004 1
F005 RS
F006 Maternal NOEL: < 3500 ppm
** Maternal NOAEL: 3500 ppm
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Pup NOEL: 3500 ppm
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     Pup NOAEL: > 7000 ppm
F007 Maternal NOEL: < 3500 ppm
* *
    Maternal NOAEL: 3500 ppm
     Pup NOEL: 3500 ppm
* *
     Pup NOAEL: > 7000 ppm
F008 HEDSET
F012 20
F020 120323
EOR
F002 27
F010 5.8.2
F004 1
F005 SO
F006 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F007 EXXON CHEMICAL, Limited Fareham, Hampshire
     EUROPEAN COMMISSION - European Chemicals Bureau Ispra (VA)
F008 CLGETTS
F009 09-01-2002
F012 20
F020 120324
EOR
F002 27
F010 5.8.2
F004 2
F005 CL
F006 Not a teratogen.
F007 Not a teratogen.
F008 CLGETTS
F012 20
F020 120325
EOR
F002 27
F010 5.8.2
F004 2
F005 RE
F006 Cox, G.E., D.E. Bailey and K. Morgareidge. (1975). Toxicity studies in
     rats with 2-butanol including growth, reproduction and teratologic
     observations. LaB No. 2093. Food and Drug Research Laboratories, Inc.,
     Waverly, NY.
F007 Cox, G.E., D.E. Bailey and K. Morgareidge. (1975). Toxicity studies in
     rats with 2-butanol including growth, reproduction and teratologic
     observations. LaB No. 2093. Food and Drug Research Laboratories, Inc.,
     Waverly, NY.
F008 CLGETTS
F012 20
F020 120326
EOR
F002 27
F010 5.8.2
F004 2
F005 RE
F006 Gallo, M.A., Oser, B.L., Cox, G.E., and Bailey, D.E. (1977). Studies on
     the long-term toxicity of 2-butanol. Toxicology and Applied Pharmacology,
     41:135 (Abstract).
F007 Gallo, M.A., Oser, B.L., Cox, G.E., and Bailey, D.E. (1977). Studies on
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the long-term toxicity of 2-butanol. Toxicology and Applied Pharmacology,
     41:135 (Abstract).
F008 CLGETTS
F012 20
F020 120327
EOR
F002 27
F010 5.8.2
F004 2
F005 RL
F006 2 - Reliable study with restrictions. No circumstances occurred that
     would have affected the quality or integrity of the data. Comparable to a
     guideline study and test procedures were in accordance with generally
     accepted scientific standa
F007 2 - Reliable study with restrictions. No circumstances occurred that
     would have affected the quality or integrity of the data. Comparable to a
     guideline study and test procedures were in accordance with generally
     accepted scientific standards and described in sufficient detail.
F008 CLGETTS
F012 20
F020 120328
EOR
F002 27
F010 5.8.2
F004 2
F005 RM
F006 Purity: > or = 99%
     Doses/Concentration: F0 Generation - second breeding.
     0, 0.3, 1.0, or 2.0% solutions (0, 538, 1644, and 3384 mg/kg-day for
    males and 0, 594, 1771, and 3122 mg/kg-day for females)
* *
    Vehicle: Drinking Water
    Number/Sex/Dose
* *
F007 Purity: > or = 99%
     Doses/Concentration: F0 Generation - second breeding.
     0, 0.3, 1.0, or 2.0% solutions (0, 538, 1644, and 3384 mg/kg-day for
    males and 0, 594, 1771, and 3122 mg/kg-day for females)
* *
    Vehicle: Drinking Water
* *
    Number/Sex/Dose:
F008 CLGETTS
F012 20
F020 120329
EOR
F002 27
F010 5.8.2
F004 2
F005 RM
F006 TERATOLOGY SCREEN RESULT: The F0 rats were mated to obtain a second
     series of pregnant dams destined to provide teratologic evaluation of the
     treatments. Pregnancy rates and survival of these females were
     unaffected. All findings sBA at bo
F007 TERATOLOGY SCREEN RESULT: The F0 rats were mated to obtain a second
     series of pregnant dams destined to provide teratologic evaluation of the
     treatments. Pregnancy rates and survival of these females were
    unaffected. All findings sBA at both 0.3 and 1.0% in the drinking water
    were negative with respect to signs of toxicity in terms of both growth
     and reproductive efficiency The body weights of the dams were not
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depressed. Examination of the uterine contents on the 20th day of

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gestation revealed that sBA was somewhat fetotoxic at the 2.0% dosage
     level, as shown by the decreased pup weights (3.74 g vs. 4.14 g in
     controls). However, that this is a minimal response is shown by the fact
     that none of the other parameters in the reproductive toxicity phase of
     this study (nidation, early or late fetal deaths) were affected. The 2.0%
    group showed apparent increases in missing sternebrae, wavy ribs, and
     incomplete vertebra ossification when compared with both the 0.3 and 1.0%
    groups. However, because the incidences for these findings in the control
    group were comparable, these effects could not be determined to be
     compound-related. The skeletal abnormalities seen in the sBA groups were
     consistent in type and frequency with the spontaneous incidence observed
     in this rat colony. There were no significant soft tissue findings in
     the 2% treated group.
F008 CLGETTS
F012 20
F020 120330
EOR
F002 27
F010 5.8.2
F004 2
F005 RS
F006 Maternal NOEL: 1% (1771 mg/kg/day)
    Maternal NOAEL: 1% (1771 mg/kg/day)
    Pup NOEL: 1% (1771 mg/kg/day)
F007 Maternal NOEL: 1% (1771 mg/kg/day)
    Maternal NOAEL: 1% (1771 mg/kg/day)
* *
     Pup NOEL: 1% (1771 mg/kg/day)
F008 CLGETTS
F012 20
F020 120331
EOB
С
Χ
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